

6 September 2002

Dear Review Committee:

Please find enclosed applications for the Critical Use Exemption from the phaseout of methyl bromide for the vegetable industry in Michigan. Specifically, we request exemption for growers of Solanaceous and Cucurbit crops who rely on fumigation for control of soilborne organisms. While my laboratory is actively researching alternative strategies, we need more time to identify and then transition the industry to viable alternatives. I did not find the proposed alternatives to methyl bromide to be scientifically proven to control *Phytophthora capsici*, a primary pathogen. Throughout the applications I've offered the research information we have regarding the proposed alternatives. Because I did not deem the proposed alternatives to be technically feasible, economic comparisons have not been included for those alternatives. Rather, we have portrayed the economics of using methyl bromide versus not using methyl bromide. It is important to note, that the economics provided do not include the economic losses during a severe disease epidemic. Just this year alone in one region of the state, we've had losses totaling over \$1 million due to this soilborne disease when methyl bromide was not used. Because of the fungicide resistance, we experienced a period from 1999 to 2000 where use of methyl bromide increased in order to manage disease.

Thank you for your attention to our applications. Please let me know if I can be of further assistance.



# DEPARTMENT OF SPLANT PATHOLOGY

Michigan State University 164 Plant Biology Building East Lansing, MI 48824-1312

> 517/353-8645 Fax: 517/353-9704

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Respectfully yours,

Mary Hausbeck
Mary Hausbeck

Professor and Extension Specialist

CUE 02 0005

For EPA Use Only ID#

# Worksheet 1. Contact and Methyl Bromide Request Information

The following information will be used to determine the amount of methyl bromide requested and the contact person for this request. It is important that we know whom to contact in case we need additional information during the review of the application.

	application.		
1.	Location (Enter the state, regi bromide.)	on, or county. Provide more detail about the	location if relevant to the feasibility of alternatives to methyl
	Michigan, USA		
2.		mmodities that benefit from the application of tween methyl bromide fumigations.)	of methyl bromide in a fumigation cycle. A fumigation cycle is
•			ucurbits include watermelon, muskmelon,
3.	cucumber, summer	squash, winter squash.	
	(Individual users sho submitting this applie	cation, please indicate the estimated percent	reviewing the U.S. climate zone map. If a consortium is tage of consortium users in each climate zone. This map is at http://www.usna.usda.gov/ Hardzone/ushzmap.html).
	All users are locate	d in zone 5B (average annual minimum te	emperature -10 to -15 F).
		-	
4.		box(es) for the soil types and percent organ se indicate the estimated percentage of con	ic matter that apply to your area. If a consortium is submitting sortium users in each soil type.
	;	Soil Type: Light X Medium	X Heavy
	Organ	ic Matter: 0 to 2%25 2 to 5 %	75 over 5%
5.	Other geographic fa	actors that may affect crop/commodity yie	eld (e.g., water table).
••	3		(
		·	
6.	Consortium name	Michigan cucurbit ( watermelon, muskmelon,	Specialty (check one)
		cucumber, summer and winter squash) growers	<u>.</u>
7.	Contact name	Dr. Mary Hausbeck	agronomic X
8.	Address	140 Plant Biology Lab	economic
		Mich. State Univ., Dept. of Plant Pathology	
		E. Lansing, MI USA 48824-1312	
9.	Daytime phone	517-355-4534	<b>10. FAX</b> 517-353-9704
11.	E-mail	hausbec1@msu.edu	
			•
	List an additional c	ontact person if available.	Specialty (check one)
12.	Contact name	Barbara Dartt, DVM, MS	agronomic
13.	Address	Salisbury Management Services, Inc.	economic X
		2487 S. Michigan, P.O. Box 10	-
		Eaton Rapids, MI USA 48827-0010	_
14.	Daytime phone	517-663-5600	- - <b>15. FAX</b> 517-663-5608
	, p		

16. E-mail

bdartt@salisbury-management.com

# Worksheet 1. Contact and Methyl Bromide Request Information

	How much active ingredient (ai) of methyl bromide are you requesting for 2005? 62,142 lbs.  If a consortium is submitting this application, the data for question 17 and 17a. should be the total for the consortium.								
In t	In the question below, area is defined as follows for each user: acres for growers, cubic feet for post harvest operations, and square feet for structural applications.								
	• •	ea will th	is be applied to? Please list units.	1,446	a	cres_units			
. Are	e vou requesting	methyl	bromide for additional years beyor	nd 2005? Yes	s X	No			
		t year and o	quantity active ingredient (ai) of methyl bro						
	Additional tin	ne is nee	ded to facilitate testing of potential	alternatives for crop sa	ifety, pathog	en			
	efficacy, and	incorpor	ation into commercial production s	systems. Also, we antic	ipate that ad	ditional			
	growing seas	ons are r	needed for demonstration plots wit	h grower cooperators.					
	If a consortium is	submitting	this application, the data below should be	the total for the consortium.					
		w, area is c	lefined as follows for each user: acres for		arvest operation	ns, and square fe			
		Year	Quantity ai (lb.) of Methyl Bromide	Area to be Treated	Unit of A	Area Treated			
		2006		1,419	8	ocres			
			60,970 a.i. (lb.)						
(Be	l-borne fungi that o	est Problible about to	58,625 a.i. (lb.)  em(s): he species or classes of pests relevant to vn, root, and fruit rot, including Phytoph	1,364 the feasibility of alternatives.)		arium			
(Be	as specific as poss	est Problible about to	58,625 a.i. (lb.)  em(s): he species or classes of pests relevant to vn, root, and fruit rot, including Phytoph	1,364 the feasibility of alternatives.)					
(Be	as specific as poss	est Problible about to	58,625 a.i. (lb.)  em(s): he species or classes of pests relevant to  vn, root, and fruit rot, including Phytoph	1,364 the feasibility of alternatives.)					
(Be	as specific as poss	est Problible about to	58,625 a.i. (lb.)  em(s): he species or classes of pests relevant to  vn, root, and fruit rot, including Phytoph	1,364 the feasibility of alternatives.)					
Soil oxy  If a issu structure.	as specific as poss  I-borne fungi that of the specific as poss  ysporum f. sp. metalogous pplying as a coruse such as size of totural applications),	est Problible about to cause crow	58,625 a.i. (lb.)  em(s): he species or classes of pests relevant to  vn, root, and fruit rot, including Phytoph	1,364 the feasibility of alternatives.)  athora capsici (primary protection)  please define a represerowers, cubic feet for post-hair	plem), and Fus	arium  Define exactly s, and square fee			
Soil oxy  If a issu structure only	pplying as a corticural applications), when pest reaches	est Problible about to cause crow donis (second means of the problem).	58,625 a.i. (lb.)  lem(s): the species or classes of pests relevant to the species of the species o	1,364 the feasibility of alternatives.) athora capsici (primary prob problem of the problem of t	entative user	arium  Define exactly s, and square fee se (treat regularly			
Soil oxy  If ap issu struu only  A re	pplying as a corress such as size of totural applications), when pest reaches	est Problible about to cause crow fonis (second meanth of the cause crow fonis fo	58,625 a.i. (lb.)  em(s): the species or classes of pests relevant to the species of	1,364 the feasibility of alternatives.) athora capsici (primary prob please define a represe rowers, cubic feet for post-han and or operation, intensity of me	entative uservest operations ethyl bromide u	arium  Define exactly and square feese (treat regularly treat regularly treats are the company to the user owns			
If a issu structure only	pplying as a corress such as size of totural applications), when pest reaches expresentative user	est Problible about to cause crow donis (second means of the cause donis of the cause done done done done done done done don	58,625 a.i. (lb.)  em(s): the species or classes of pests relevant to the species of	the feasibility of alternatives.)  athora capsici (primary protein pro	entative user rvest operations ethyl bromide u	arium  Define exactly and square feese (treat regularly treat regularly treats are the company to the user owns			
If a issu structure only	pplying as a corress such as size of totural applications), when pest reaches expresentative user	est Problible about to cause crow donis (second means of the cause donis of the cause done done done done done done done don	58,625 a.i. (lb.)  lem(s): the species or classes of pests relevant to the species of the species o	the feasibility of alternatives.)  athora capsici (primary protein pro	entative user rvest operations ethyl bromide u	arium  Define exactly and square feese (treat regularly treat regularly treats are the company to the user owns			
If a issu structure only  A return for t	pplying as a corress such as size of totural applications), when pest reaches expresentative user land and operation the fresh market in	est Problible about to cause crow donis (secondary) and the operation whether the cauth at threshold employs reductive, and dustry, and adustry, and the operation of the operat	58,625 a.i. (lb.)  lem(s): the species or classes of pests relevant to the species of the species o	the feasibility of alternatives.)  athora capsici (primary protein pro	entative user rvest operations ethyl bromide u	arium  Define exactly and square feese (treat regularly treat regularly treats are the company to the user owns			
If applied the fort	pplying as a corress such as size of totural applications), when pest reaches expresentative user land and operation the fresh market in a. Explain why the	est Problible about to cause crow donis (secondary) and the operation whether the cause at threshold employs readustry, and this user to the operation of the user t	58,625 a.i. (lb.)  lem(s): the species or classes of pests relevant to the species of	the feasibility of alternatives.)  athora capsici (primary protein pro	entative user rvest operations ethyl bromide u olied preplant. ated regularly. stharvest.	arium  Define exactly and square feese (treat regularly treat regularly the user owns).			

# Worksheet 2-A. Methyl Bromide - Use 1997-2000

Col A: Formulation of Methyl Bromide	averages for proportions	Enter the appropriate data in Col B-M for each formulation, if known, and/or the totals and averages for all formulations. If you enter only the total and averages for all formulations in the last row of the table, please describe in the comments section the formulations typically used, or the approximate proportions of the formulations used.										
Col B, E, H, K: Actual Area Treated	consortium	, for the year	indicated.							otal actual are		e 
	individual u	iser or the en	tire consortiur	n, for the year	indicated.					ounds ai applie	d by the	
Col D, G, J, M: Actual Average lbs. ai Applied per Area							omatically cald		ne previous 2	columns.		
Area is defined below as follows for each to	user: acres for g	rowers, cubic	feet for post-	harvest opera	tions, and squ	are feet for s	tructural appli	cations.		<del></del>		<del></del>
Α	В	С	D	Е	F	G	Н		J	K	L	M
Formulation of Methyl Bromide		1997			1998			1999			2000	
	Total Actual Area Treated	Actual Total Ibs. ai of Methyl Bromide Applied	Average Ibs. ai Applied per Area	Total Actual Area Treated	Actual Total lbs. ai of Methyl Bromide Applied	Average Ibs. ai Applied per Area	Total Actual Area Treated	Actual Total lbs. ai of Methyl Bromide Applied	Average Ibs. ai Applied per Area	Total Actual Area Treated	Actual Total Ibs. ai of Methyl Bromide Applied	Averag Ibs. a Applied Area
over 95% methyl bromide										<b>!</b>		
over 95% methyl bromide 75% methyl bromide, 25% chloropicrin												
•	568	60970	107.341549	546	58625	107.371795	546	58625	107.371795	622	66833	107.448
75% methyl bromide, 25% chloropicrin	568	60970	107.341549	546	58625	107.371795	546	58625	107.371795	622	66833	107.448
75% methyl bromide, 25% chloropicrin 67% methyl bromide, 33% chloropicrin	568	60970	107.341549	546	58625	107.371795	546	58625	107.371795	622	66833	107.448
75% methyl bromide, 25% chloropicrin 67% methyl bromide, 33% chloropicrin 50% methyl bromide, 50% chloropicrin	568	60970	107.341549	546	58625	107.371795	546	58625	107.371795	622	66833	107.448

Comments:

Actual area treated is in acres. Applications were made under plastic with an average bed width of 24".

Application for Critical Use Exemption of Methyl Bromide for Use in 2005 in the United States

### **Economic Summary:**

Budgets representative of eight Michigan fresh market vegetable crops were constructed using grower focus groups. These budgets were compared to the alternative revenue and cost structures present if no methyl bromide were utilized. In the case of all eight crops, use of methyl bromide generates higher profits than production without this fumigant.

## **Application for Critical Use Exemption of Methyl Bromide** for Use in 2005 in the United States

## Worksheet 2-D. Methyl Bromide - Use and Costs for 2001

In Michigan fresh market vegetable crops, methyl bromide (MB) is utilized as a component of a plasticulture system. Fields are prepared with a minimal amount of traditional tillage using moldboard plows, tandem discs and tractors of 100-140 horsepower. Following this preparation, a tractor (100 hp) pulls a piece of equipment called a bedder or plastic layer. This equipment simultaneously forms beds, lays plastic and drip tube and injects MB. This equipment can usually bed about 1 acre per hour and utilizes a crew of 4 laborers and an equipment operator. The operating and ownership costs of running this equipment and paying the crew were calculated to be \$98 per acre. It is not possible to separate the application of MB from the portions of the plasticulture system.

## Worksheet 2-C. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 2001

Eight vegetable crops are included in this application. They are:

Curcubit Crops

Solanaceous Crops

- Cantaloupe
- Eggplant
- Cucumber

- Tomato
- Hard Squash

- Green Pepper
- Watermelon
- Zucchini

Price data for these crops were averaged across quality grades and seasonal differences. Use of MB does facilitate earlier harvests because it has a shorter spring waiting period than alternatives. Methyl bromide also supports growth of a healthier, more vigorous plant. On average, crops produced with methyl bromide bring a higher price because they can be marketed in more timely manner, taking advantage of early high prices. In addition, a larger percentage of product can be marketed in quality and size grades that bring a higher price.

## Worksheet 2-E. Methyl Bromide - Other Operating Costs for 2001

## Methodology and Assumptions for Budgets Provided in Place of Worksheet 2-E.

The budgets for the eight crops included in this application were developed using grower focus groups with a good knowledge of the industry and good field, enterprise, and financial records. The process was initiated by defining individual production systems representative of Michigan. Subsequently, both the sequence of decisions and the information necessary to make these key decisions was collected. This process resulted in a list of inputs and input prices that were then translated into costs. These costs were verified against grower records. These budgets reproduce, as completely as possible, all costs incurred by growers.

Below are comments about the methods used in particular areas of each budget.

## Costs of Capital Services (Buildings, Machinery, and Equipment)

Estimating the annual cost of using buildings, machinery, equipment and other assets is a challenge in cost of production studies. Buildings, machinery and services were priced to the enterprise on a "custom" basis. Further, services such as land preparation were priced to the enterprise as a "bundled" service/task reflecting both the machinery and labor components of the service.

This approach requires some judgment because costs such as buildings to house machinery and equipment, the farm shop, and labor used in maintenance of machinery and equipment must be included in the "custom fee" as well as the "depreciation and interest" on the machinery and equipment. The fact that this custom fee approach was used does not imply that custom operators did all the tasks. It simply means the tasks are priced to the enterprise as if a custom operator had completed them. The services may well have been provided by the "machinery services enterprise" of the farm. As a double check, members of the focus group attempted to compare the aggregate custom fee costs to those based on their accounting records which included labor, custom fees, and depreciation and interest on buildings, machinery, and equipment. Custom fees were also double-checked against survey information when available.

# Worksheet 2-F. Methyl Bromide Fixed and Overhead Costs in 2001

Fixed costs including management and supervision, insurance and other overhead were allocated equally across an entity's total vegetable acres. In the case of management and labor, adjustments were made to account for increased time demands of crops with a more complex biological or production cyce.

ID#

# Worksheet 2-B. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 1997-2000

If a same ordination			proofiffically 11	elu allu Gloss F	kevenue 1997-2000		
The purpose of the	submitting this	application, the data for t	his table should reflect the	actual averages for the	consortium.		
Cal At Va-	The purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify this  Col. A: Year  Be sure to enter the year. Use as many rows as needed for each year for all the crops/commodities in the fumigation cycles from 1997 to						
Col. B: Crop/Con	nmodity	Citici all cropa/collin	iodines mai benent nom m	leinvi bromide in each fui	migation cycle /For example #		
Col. B: Crop/Commodity  Enter all crops/commodities that benefit from methyl bromide in each fumigation cycle. (For example, if normally methyl bromide is ap  Col. C: Unit of  Enter all crops/commodities that benefit from the application of methyl bromide in the fumigation cycle and you do not have the						t you do not have the	
Col. C: Unit of		Lines are unit of frea	surement for each crop/cc	mmodity.	2 and ramigation by old and	you do not have the	
Col. D: Crop/Con	nmodity Yield	Enter the number of a	units of crop/commodities	produced per area.			
Col. E: Price		Enter the average pri	ces received by the users	for the year and cron/cor	mmodity indicated (1997-2000).		
Col. F: Revenue		This number is calcul	ated automatically using the	ne values you entered in	Cols. D and E. You may override the	formula to automorphic	
Total Revenue for	r 1997-2000	Enter the total revenu	e per year by adding the r	evenue for all crops for the	hat year	iornula to enter a different	
Average Revenue	per Year:	The average revenue	per vear is calculated aut	omatically using the sum	many data you anter fee		
Area is defined be	elow as follows	for each user: acres for	growers, cubic feet for pos	t-harvest operations, and	mary data you enter for each year.  I square feet for structural applications	·	
Α		В	C	D	square feet for structural applications		
Year	Cro	p/Commodity	Unit of	Crop/Commodity		F	
Methyl Bromide		•	Crop/Commodity	Yield	Price	Revenue	
was Applied			(e.g., pounds, bushels)	(Units per area)	(per unit of crop/commodity)	(per area)	
1997	C	Canteloupe	700 lb bin				
1998	Canteloupe		700 lb bin	53 53	\$ 125.00	\$ 6,625.00	
	Canteloupe		700 lb bin	53	\$ 120.00	\$ 6,360.00	
2000	Canteloupe		700 lb bin		\$ 110.00	\$ 5,830.00	
			1 00 15 511	53	\$ 90.00	\$ 4,770.00	
				.p		#VALUE!	
						\$ 0.00	
			1			\$ 0.00	
						\$ 0.00	
						\$ 0.00	
						\$ 0.00	
						\$ 0.00	
			······································		Total Revenue for 1997	\$ 0.00	
				}-	Total Revenue for 1998	\$ 6,625.00	
				-	Total Revenue for 1999	\$ 6,360.00	
				-	Total Revenue for 1999	\$ 5,830.00	
				}-	Average Revenue Per Year	\$ 4,770.00	
						\$ 5,896.25	



# Michigan Cantalouple WITH Methyl Bromide

Price Quantity Unit	per Unit	Total per Acre
Revenue		
Cantaloupe 57 700 lb bin \$ 90.	00 \$	5,130
Total Revenue	\$	5,130
Expenses		
Field Preparation		
Cover Crop (Materials, Machinery & Labor)	\$	13
Soil Test	. \$	2
Lime & Application	\$	13
Fertilizer (Materials)	\$	100
Apply & Incorporate Fertilizer (Machinery & Labor)	\$	20
Plastic & Drip Tape (Materials)	\$	284
Lay Plastic & Drip Tape (Machinery & Labor)	\$	. 98
Fumigate (Materials)	\$	341
Herbicide, Insecticide & Fungicide (Materials)	\$	183
Apply Herbicide, Insecticide & Fungicide (Mach & Lab)	\$	78
Plant & Grow	·	
Transplants	\$	435
Planting (Machinery & Labor)	\$	65
Cultivate, Hoe & Move Vines (Mach & Lab)	\$	108
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	\$	207
Pollination (Bees)	\$	35
Scouting & Lab Work	\$	30
Field Maintenance - Driveways & Mowing (Mach & Lab)		7
Harvest		
Pick Crop (Field to Packing Shed)	\$	855
Grading & Packing (Includes shipping containers)	\$	713
Shipping (Includes Materials, Machinery & Labor)	\$	71
Sales & Marketing 9% of gross	\$	462
Field Clean-up	\$	112
Management & Supervision	\$	75
Interest on Operating Capital 8%	\$	103
Land Rent	\$	200
Insurance	\$	8
Other Overhead (Professional Fees, Education & Travel, etc)	\$	10
Total Expenses	\$	4,626
PROFIT	\$	504

# Michigan Cantalouple WITHOUT Methyl Bromide

Price		Total per	
Quantity Unit	Unit	Acre	
Revenue			
Cantaloupe 34 700 lb bin \$ 85.	.00 \$	2,907	
Total Revenue	\$	2,907	
Expenses			
Field Preparation			
Cover Crop (Materials, Machinery & Labor)	\$	13	
Soil Test	\$	2	
Lime & Application	\$	13	
Fertilizer (Materials)	\$	100	
Apply & Incorporate Fertilizer (Machinery & Labor)	\$ \$		
Plastic & Drip Tape (Materials)	э \$	20	
Lay Plastic & Drip Tape (Machinery & Labor)	\$ \$	284	
Fumigate (Materials)	Φ	98	
Herbicide, Insecticide & Fungicide (Materials)	·	050	
Apply Herbicide, Insecticide & Fungicide (Mach & Lab)	\$ \$	253	
Plant & Grow	Ф	90	
Transplants	œ	435	
Planting (Machinery & Labor)	\$ \$		
Cultivate, Hoe & Move Vines (Mach & Lab)	\$ \$	65 143	
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	э \$	143	
Pollination (Bees)	э \$	207	
Scouting & Lab Work	э \$	35 30	
Field Maintenance - Driveways & Mowing (Mach & Lab)	\$ \$	30	
Harvest	Ф	7	
Pick Crop (Field to Packing Shed)	•	540	
Grading & Packing (Includes shipping containers)	\$	513 428	·
Shipping (Includes Materials, Machinery & Labor)	\$	420	
O-1 0 B4- 1 1'	\$	43	
Sales & Marketing 9% of gross Field Clean-up	\$	262	
Management & Supervision	\$	112	
	\$	75 50	
Interest on Operating Capital 8% Land Rent	\$	58	
Insurance	\$	200	
· · · · · · · · · · · · · · · · · · ·	\$	8	
Other Overhead (Professional Fees, Education & Travel, etc)	\$	10	
Total Expenses	\$	3,502	
PROFIT	\$	(595)	

# Worksheet 2-B. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 1997-2000

If a consortium is submitting this ap	oplication, the data for this table should reflect the actual averages for the consortium.
	estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify this in operations when providing gross revenue data.
Col. A: Year	Be sure to enter the year. Use as many rows as needed for each year for all the crops/commodities in the fumigation cycles from 1997 to 2000. If a fumigation cycle overlaps more than one calendar year, then the year of the fumigation cycle is the year methyl bromide was applied.
Col. B: Crop/Commodity	Enter all crops/commodities that benefit from methyl bromide in each fumigation cycle. (For example, if normally methyl bromide is applied and tomatoes are grown and harvested followed by peppers without an additional treatment of methyl bromide, then both tomatoes and peppers would be part of the same fumigation cycle.) See the Fumigation Cycle Worksheet for a comprehensive definition of the fumigation cycle.
	If someone other than the applicant benefits from the application of methyl bromide in the fumigation cycle and you do not have the quantitative data for the crops grown on the same land, please indicate so in the comments section below.
Col. C: Unit of Crop/Commodity	Enter the unit of measurement for each crop/commodity.
Col. D: Crop/Commodity Yield	Enter the number of units of crop/commodities produced per area.
Col. E: Price	Enter the average prices received by the users for the year and crop/commodity indicated (1997-2000).
Col. F: Revenue	This number is calculated automatically using the values you entered in Cols. D and E. You may override the formula to enter a different revenue. Please explain why the revenue amount is different in the comment section below.
Total Revenue for 1997-2000	Enter the total revenue per year by adding the revenue for all crops for that year.
Average Revenue per Year:	The average revenue per year is calculated automatically using the summary data you enter for each year.
Area is defined below as follows f	or each user: acres for growers, cubic feet for post-harvest operations, and square feet for structural applications.

А	В	С	D	E	F
Year Methyl Bromide was Applied	Crop/Commodity	Unit of Crop/Commodity (e.g., pounds, bushels)	Crop/Commodity Yield (Units per area)	Price (per unit of crop/commodity)	Revenue (per area)
1997	Cucumber	1 1/9 Bushel boxes	750	\$ 11.31	\$ 8,480.77
1998	Cucumber	1 1/9 Bushel boxes	800	\$ 13.02	\$ 10,417.58
1999	Cucumber	1 1/9 Bushel boxes	825	\$ 9.79	\$ 8,077.75
2000	Cucumber	1 1/9 Bushel boxes	900	\$ 10.82	\$ 9,741.76
					\$ 0.00
					\$ 0.00
	·				\$ 0.00
				··	\$ 0.00
	<u></u>		<del></del>		\$ 0.00
ļ.,		-			\$ 0.00
	<u> </u>				\$ 0.00
1				Total Revenue for 1997	\$ 8,480.77
			<u> </u>	Total Revenue for 1998	\$ 10,417.58
			1	Total Revenue for 1999	\$ 8,077.75
Ì			1	Total Revenue for 2000	\$ 9,741.76
ļ			L	Average Revenue Per Year	\$ 9,179.46

Comments:



# Michigan Cucumber WITH Methyl Bromide

Quantity U	nit	Price per Unit		Total per Acre
Revenue	•	10.00	Φ	10 505
Cucumber 1,025 boxe	\$	12.22	\$	12,525
Total Revenue			\$	12,525
Expenses				
Field Preparation				
Cover Crop (Materials, Machinery & Labor)			\$	13
Soil Test			\$	2
Lime & Application			\$	13
Fertilizer (Materials)			\$	100
Apply & Incorporate Fertilizer (Machinery & L	.abor)		\$	20
Plastic & Drip Tape (Materials)			\$	284
Lay Plastic & Drip Tape (Machinery & Lab	or)		\$	98
Fumigate (Materials)	·		\$	341
Herbicide, Insecticide & Fungicide (Material	s)		\$	183
Apply Herbicide, Insecticide & Fungicide (Mach	& Lab	)	\$	54
Plant & Grow				
Transplants			\$	813
Planting (Machinery & Labor)			\$	65
Cultivate, Hoe & Move Vines (Mach & Lab)			\$	108
Drip Irrigate & Fertigate (Materials, Labor & El	ectrici	ty)	\$	207
Pollination (Bees)			\$	35
Scouting & Lab Work			\$	30
Field Maintenance - Driveways & Mowing	(Macl	n & Lab)	\$	7
Harvest				
Pick Crop (Field to Packing Shed)			\$	1,230
Grading & Packing (Includes shipping containe	s)		\$	3,065
Shipping (Includes Materials, Machinery & Labor)			\$	513
Field Clean-up			\$ \$ \$	112
Sales & Marketing 9% of gro	ss		\$	1,127
Management & Supervision			\$	75
Interest on Operating Capital 8	%		\$	251
Land Rent	•		\$	200
Insurance		٠.	\$	8
Other Overhead (Professional Fees, Education & T	ravel	, etc)	\$	10
Total Expenses		<del></del>	\$	8,963
PROFIT			\$	3,562

# Michigan Cucumber WITHOUT Methyl Bromide

Price pe Quantity Unit Un		Total per Acre
Revenue		
Cucumber 615.0 11/9 bu boxes \$ 10.22	\$	6,285
Total Revenue	\$_	6,285
Expenses		
Field Preparation		
Cover Crop (Materials, Machinery & Labor)	\$	13
Soil Test	\$	2
Lime & Application		13
Fertilizer (Materials)	\$	100
Apply & Incorporate Fertilizer (Machinery & Labor)	\$ \$ \$	20
Plastic & Drip Tape (Materials)	\$	284
Lay Plastic & Drip Tape (Machinery & Labor)	\$	98
Fumigate (Materials)		
Herbicide, Insecticide & Fungicide (Materials)	\$	253
Apply Herbicide, Insecticide & Fungicide (Mach & Lab)	\$	54
Plant & Grow		
Transplants	\$	813
Planting (Machinery & Labor)	\$	65
Cultivate, Hoe & Move Vines (Mach & Lab)	\$	143
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	\$	207
Pollination (Bees)	\$	35
Scouting & Lab Work	\$	30
Field Maintenance - Driveways & Mowing (Mach & Lab)	\$	7
Harvest	_	
Pick Crop (Field to Packing Shed)	\$	738
Grading & Packing (Includes shipping containers)	\$	1,839
Shipping (Includes Materials, Machinery & Labor)	\$	308
Field Clean-up	\$	112
Sales & Marketing 9% of gross	\$	566
Management & Supervision	<b>\$</b>	75
Interest on Operating Capital 8%	Þ	126
Land Rent	\$ \$ \$ \$ \$	200
Insurance Other Overhead (Professional Fees, Education & Travel, etc)	\$ \$	8 10
	φ	10
Total Expenses	\$	6,117
PROFIT	\$	168

# Worksheet 2-B. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 1997-2000

AAOLV2LICEL T-r	J. Michigi Diolingo Cici							
If a consortium is su	ubmitting this application, the data for th	is table should reflect the a	actual averages for the c	consortium.	and with EDA to modify this			
The purpose of this	f a consortium is submitting this application, the data for this table should reflect the actual averages for the consortium.  The purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify the purpose of this worksheet is to estimate the gross revenue for 1997 to the purpose of the growth and the growth							
Col. A: Year								
	57 t	adition that banefit from me	sthyl bromide in each fum	igation cycle. (For example, if normal	is theftist profitible is abblica			
Col. B: Crop/Com	modity Effet all crops/comm	the applicant benefits from	n the application of meth	yl bromide in the fumigation cycle and	you do not have the			
	it someone other trial	Title applicant benefits from	nmodity					
Col. C: Unit of	Enter the unit of mea	surement for each crop/cor	minouity.					
Col. D: Crop/Com	modity Yield Enter the number of u	inits of crop/commodities p	roduced per area.	dity indicated (1007 2000)				
Col. E: Price		ces received by the users f	or the year and crop/com	modity indicated (1997-2000).	formula to enter a different			
Col. F: Revenue	This number is calcul	ated automatically using th	e values you entered in C	Cols. D and E. You may overlide the t	officia to effer a different			
Total Revenue for	Catas the total revenu	e per year by adding the re	evenue for all crops for th	at year.				
	The everage revenue	ner vear is calculated auto	omatically using the sumr	hary data you enter for each year.				
Average Revenue	low as follows for each user: acres for	groupes cubic feet for nos	t-harvest operations, and	square feet for structural applications				
Area is defined be	low as follows for each user: acres for	growers, cabic reet for posi-	D.	F	F			
Α	В	C	<u> </u>	Price	Revenue			
7.6	Cron/Commodity	Unit of	Crop/Commodity	Price	Nevende			

	B	С	D	E '	F
Year Methyl Bromide was Applied	Crop/Commodity	Unit of Crop/Commodity (e.g., pounds, bushels)	Crop/Commodity Yield (Units per area)	Price (per unit of crop/commodity)	Revenue (per area)
	Hard Sayach	Bushel (60 lbs.)	300	\$ 6.54	\$ 1,963.
	Hard Squash	Bushel (60 lbs.)	300	\$ 6.54	\$ 1,963
	Hard Squash	Bushel (60 lbs.)	500	\$ 8.74	\$ 4,370
	Hard Squash	Bushel (60 lbs.)	500	\$ 11.90	\$ 5,947
2000	Hard Squash	Dustier (oo ibs.)			\$ 0
					\$ (
					\$ (
					\$ (
					\$ (
					\$ (
					\$ (
					\$ (
				Total Revenue for 1997	\$ 1,963
			<u> </u> -	Total Revenue for 1998	\$ 1,963
	·		<b>}</b>	Total Revenue for 1999	\$ 4,37
			<u> </u>	Total Revenue for 2000	\$ 5,94
			f	Average Revenue Per Year	\$ 3,56

Comments:



# Michigan Winter Squash WITH Methyl Bromide

Price pe Quantity Unit Un		Total per Acre				
Revenue						
Winter Squash 500 bu \$ 10.34	\$	5,168				
	•	,				
Total Revenue	\$_	5,168				
Expenses						
Field Preparation						
Cover Crop (Materials, Machinery & Labor)	\$	13				
Field Tillage (Moldboard plow, Disc twice)	\$	39				
Soil Test	\$ \$ \$ \$ \$ \$ \$ \$ \$	2				
Lime & Application	\$	13				
Fertilizer (Materials)	\$	75				
Apply & Incorporate Fertilizer (Machinery & Labor)	\$	20				
Plastic & Drip Tape (Materials)	\$	284				
Lay Plastic & Drip Tape (Machinery & Labor)	\$	98				
Fumigate (Materials)	\$	341				
Herbicide, Insecticide & Fungicide (Materials)		325				
Apply Herbicide, Insecticide & Fungicide (Mach & Lab) \$						
Plant & Grow	_					
Transplants	\$	442				
Planting (Machinery & Labor)	\$	65				
Cultivate, Hoe & Move Vines (Mach & Lab)	\$ \$ \$	53				
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	\$	207				
Pollination (Bees)	\$	35				
Scouting & Lab Work	\$	30				
Field Maintenance - Driveways & Mowing (Mach & Lab)	\$	7				
Harvest	•	450				
Pick Crop (Field to Packing Shed)	\$	450 1,145				
Grading & Packing (Includes shipping containers)	\$	1,145 250				
Shipping (Includes Materials, Machinery & Labor)	\$ \$ \$ \$ \$ \$ \$	465				
Sales & Marketing 9% of gross	φ	112				
Field Clean-up	Φ	70				
Management & Supervision Interest on Operating Capital 8%	φ Φ	103				
miorodi or operaning a special	Ψ	200				
Land Rent	\$	8				
Insurance Other Overhead (Professional Fees, Education & Travel, etc)	Ф \$	10				
Other Overhead (Floressional Fees, Education & Havet, etc)	Ψ	10				
Total Expenses	\$	4,917				
PROFIT	\$	251				

# Michigan Winter Squash WITHOUT Methyl Bromide

Price Quantity Unit	per Unit	Total per Acre	
Revenue			
Winter Squash 375 bu \$ 7.	.09 \$	2,660	
Fotal Revenue	\$	2,660	
Expenses			
Field Preparation			
Cover Crop (Materials, Machinery & Labor)	\$	13	
Field Tillage (Moldboard plow, Disc twice)	\$	39	· : - %:
Soil Test	\$	2	
Lime & Application	\$	13	
Fertilizer (Materials)	\$	75	
Apply & Incorporate Fertilizer (Machinery & Labor)	\$ \$ \$	20	
Plastic & Drip Tape (Materials)	\$	284	
Lay Plastic & Drip Tape (Machinery & Labor)	\$	98	
Fumigate (Materials)	,	- <del>-</del>	
Herbicide, Insecticide & Fungicide (Materials)	\$	403	
Apply Herbicide, Insecticide & Fungicide (Mach & Lab)	\$	60	
Plant & Grow	7		
Transplants	\$	442	
Planting (Machinery & Labor)	\$	65	
Cultivate, Hoe & Move Vines (Mach & Lab)	\$	68	
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	\$	207	
Pollination (Bees)	\$ \$ \$	35	
Scouting & Lab Work	\$	30	
Field Maintenance - Driveways & Mowing (Mach & Lab		7	
Harvest	, +	•	
Pick Crop (Field to Packing Shed)	\$	338	
Grading & Packing (Includes shipping containers)	\$	859	
Shipping (Includes Materials, Machinery & Labor)	\$	188	
Sales & Marketing 9% of gross	\$	239	
Field Clean-up	\$	112	
Management & Supervision	\$	70	
Interest on Operating Capital 8%	\$	53	
Land Rent	\$	200	
Insurance	\$	8	
Other Overhead (Professional Fees, Education & Travel, etc)	э \$	0 10	
		10	
Total Expenses	\$	3,938	
PROFIT	\$	(1,278)	

# Worksheet 2-B. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 1997-2000

	~. inctill	y: Diviliae • 6(6	NA AIIDOUIIIOONA	eld and Groce !	20V00110 4007 2002	
If a consortium is submitting this application, the data for this table should reflect the actual averages for the consortium.						
The purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify this  Col. A: Year  Be sure to enter the year. Use as many rows as needed for each year for all the except year for all the except year.						
Col. A: Year  Be sure to enter the year. Use as many rows as needed for each year for all the crops/commodities in the fumigation cycles from 1997 to  Col. B: Crop/Commodity  Enter all crops/commodities that benefit from methyl bromide in each fumigation cycles from 1997 to					work with EPA to modify this	
Col. B: Crop/Commodity  Enter all crops/commodities that benefit from methyl bromide in each if someone other than the applicant benefits from the application of				s needed for each year f	or all the crops/commodities in the fum	igation cycles from 1997 to
		If someone other tha	in the applicant benefits for	lethyl bromide in each fu	migation cycle. (For example, if norma	ally methyl bromide is applied
Col. C: Unit of	<del></del>	Enter the unit of mea	asurement for each crop/co	m the application of met	migation cycle. (For example, if norma hyl bromide in the fumigation cycle and	you do not have the
Col. D: Crop/Cor	nmodity Yield		units of crop/commodities			
Col. E: Price		Enter the average or	ices received by the	produced per area.		
Col. F: Revenue	······································	This number is calcu	lated automatically are users	for the year and crop/co	mmodity indicated (1997-2000).	
Total Revenue for		Time marrison to dated	idica automatically using tr	16 Values vou entered in	Cole Dand E. Vou man avanish it	formula to enter a different
Average Revenue	ner Year	The average rovenus	ue per year by adding the r	evenue for all crops for t	hat year.	
Area is defined be	elow as follows	for each upor: acros for	per year is calculated auto	omatically using the sum	nat year. mary data you enter for each year.	
A	0.011 0.0101013	B	growers, cubic feet for pos	t-harvest operations, and	mary data you enter for each year.  I square feet for structural applications	
Year		p/Commodity	<u> </u>	D	E	F
Methyl Bromide	Cit	prominiouity	Unit of	Crop/Commodity	Price	Revenue
was Applied			Crop/Commodity	Yield	(per unit of crop/commodity)	(per area)
	Watermelon - 8	Conded	(e.g., pounds, bushels)	(Units per area)		(por dica)
1997	Watermelon - S	Seeded	pound	12,000	\$ 0.09	\$ 1,080.00
	Watermelon - 8		pound	42,000	\$ 0.15	\$ 6,090.00
1990	Watermelon - S	Seeded	pound	12,000	\$ 0.10	
1990	Watermelon - S	Seedless	pound	42,000	\$ 0.13	\$ 1,200.00
1999	Watermelon - S	· · · · · · · · · · · · · · · · · · ·	pound	12,000	\$ 0.10	\$ 5,460.00
2000	Watermelon - S		pound	42,000	\$ 0.13	\$ 1,140.00 \$ 5,460.00
2000	Watermelon - S	·	pound	12,000	\$ 0.11	\$ 1,320.00
2000	watermeion - S	seedless	pound	42,000	\$ 0.11	\$ 4,620.00
						\$ 4,620.00
	<del></del>					\$ 0.00
						\$ 0.00
——————————————————————————————————————		<del></del>				\$ 0.00
					Total Revenue for 1997	\$ 7,170.00
				1	Total Revenue for 1998	\$ 6,660.00
					Total Revenue for 1999	\$ 6,600.00
					Total Revenue for 2000	\$ 5,940.00
Average Davis B. V.					\$ 6,592.50	
				_		Ψ 0,032.30
						i
· · · · · · · · · · · · · · · · · · ·						ſ



# Michigan Watermelon WITH Methyl Bromide

	With Methyl B	CHIR	1 <del>C</del>		
			Price pe	r	Total per
	Quantity	Unit	Uni	t	Acre
Revenue					
Seedless	42,000	lb S	-	\$	5,460
Seeded	12,000	lb S	6 0.12	\$	1,440
Total Revenue	·			\$	6,900
Expenses					•
Field Preparation					
Cover Crop (Mat	erials, Machinery & Labo	r)		\$	13
Field Tillage (Mo	ldboard plow, Disc twice)	)		\$	39
Soil Test				\$	2
Lime & Applicat	ion			\$	13
Fertilizer (Materia				\$ \$	121
	rate Fertilizer (Machir	ery & La	abor)	\$	20
Plastic & Drip Ta				\$	284
	ip Tape (Machinery	& Labo	or)	\$.	98
Fumigate (Materia	als)		·	\$	341
Herbicide, Insec	ticide & Fungicide (M	<b>Materials</b>	;)	\$	280
Apply Herbicide, I	nsecticide & Fungicide	(Mach 8	Lab)	\$	48
Plant & Grow					
Seeds (Seedless &	k Seeded)			\$	173 .
Growing Seeds			•	\$	144
Planting				\$ \$	65
Cultivate, Hoe &	Move Vines (Mach &	Lab)		\$	62
Drip Irrigate & F	ertigate (Materials, Lab	or & Ele	ctricity)	\$	207
Pollination (Bees	s)			\$	54
Scouting & Lab				\$	30
Field Maintenan	ce - Driveways & Mo	owing (	Mach & Lab	)	7
Harvest					
Pick Crop (Field to	Packing Shed)			\$	900
Grading & Packi	ng (Includes shipping co	ontainers	5)	\$	300
Shipping (Includes	Materials, Machinery &	Labor)		\$	75
Sales & Marketing	9% of	gross		\$	621
Field Clean-up				\$	112
Management & Su	pervision (Includes pic	kup)		\$ \$ \$ \$ \$ \$	85
Interest on Operat	ing Capital	8%		\$	138
Land Rent		200			
Insurance				\$	8
Other Overhead (P	rofessional Fees, Educat	ion & Tr	avel, etc)	\$	10
Total Expenses				\$	4,450
PROFIT				\$	2,450

# Michigan Watermelon WITHOUT Methyl Bromide

	Ouantity	Unit	Price per Unit		Total pe
Revenue	Quantity	Unit	Uni		Acr
Seedless	29,400	lb \$	0.13	\$	2 022
Seeded	8,400	lb \$		Ф \$	3,822
Occueu	0,400	υφ	0.12	Ф	1,008
Total Revenue				\$	4,830
Expenses					
Field Preparation					
Cover Crop (Materials, M	achinery & Labo	or)		\$	13
Field Tillage (Moldboard	plow, Disc twice	)		\$	39
Soil Test				\$	2
Lime & Application				\$	13
Fertilizer (Materials)				\$	121
Apply & Incorporate Fe	rtilizer (Machi	nery & Lab	or)	\$	20
Plastic & Drip Tape (M	aterials)			\$	284
Lay Plastic & Drip Tape	e (Machinery	& Labor	·)	\$	98
Fumigate (Materials)	1				
Herbicide, Insecticide 8	ß Fungicide (	Materials)		\$	405
Apply Herbicide, Insectici	de & Fungicid	e (Mach & L	₋ab)	\$	66
Plant & Grow				•	
Seeds (Seedless & Seeder	d)			\$	173
Growing Seeds				\$	144
Planting				\$	65
Cultivate, Hoe & Move	Vines (Mach 8	k Lab)		\$	162
Drip Irrigate & Fertigate	(Materials, Lat	or & Elect	ricity)	\$	207
Pollination (Bees)				\$	54
Scouting & Lab Work				\$	30
Field Maintenance - Dr	iveways & M	owing (M	ach & Lab)	•	
Harvest	,	•	•		
Pick Crop (Field to Packin	g Shed)			\$	630
Grading & Packing (Incl	udes shipping c	ontainers)		\$	210
Packing Shed, Equip &		,		\$	53
Sales & Marketing	9% d	of gross		\$	435
Field Clean-up		<b>5</b>		\$	112
Management & Supervisi	ON (Includes pi	ckup)		\$	85
nterest on Operating Car		8%		\$ \$ \$	97
and Rent				\$	200
nsurance				\$	8
Other Overhead (Profession	nal Fees, Educa	tion & Tra	vel, etc)	\$	10
Total Expenses				\$	3,742
PROFIT				\$	1,088

### Worksheet 2-B. Methyl Bromide - Crop/Commodity Yield and Gross Revenue 1997-2000

VOIRSITEEL 2-			•			
The purpose of the	submitting this a	ipplication, the data for the	his table should reflect the	actual averages for the	consortium.	
The purpose of this worksheet is to estimate the gross revenue for 1997 - 2000 when using methyl bromide. Post-harvest and structural users may work with EPA to modify this						
Col. A: Year  Be sure to enter the year. Use as many rows as needed for each year for all the crops/commodities in the fumigation cycles from 1997 to  Col. B: Crop/Commodity  Enter all crops/commodities that benefit from methyl bromide in each fumigation cycle. (For example, if normally methyl bromide is applie						
Col. B: Crop/Con	nmodity	Enter all crops/comm	lodities that benefit from me	ethyl bromide in each fur	nigation cycle. (For example, if norma	lly methyl bromide is applied
Col. C: Unit of					nyl bromide in the fumigation cycle and	you do not have the
			surement for each crop/cor			
Col. D: Crop/Con	nmodity Yield		units of crop/commodities p			
Col. E: Price					nmodity indicated (1997-2000).	
Col. F: Revenue					Cols. D and E. You may override the	formula to enter a different
Total Revenue for			ue per year by adding the re			
Average Revenue					mary data you enter for each year.	
	elow as follows	for each user: acres for			square feet for structural applications	
A			C	D	E	F
Year	Crop	p/Commodity	Unit of	Crop/Commodity	Price	Revenue
Methyl Bromide			Crop/Commodity	Yield	(per unit of crop/commodity)	(per area)
was Applied			(e.g., pounds, bushels)	(Units per area)		
	Zucchini		1/2 Bushel (20 lbs.)	750	\$ 4.62	
	Zucchini		1/2 Bushel (20 lbs.)	750	\$ 4.62	\$ 3,465.66
	Zucchini		1/2 Bushel (20 lbs.)	1000	\$ 6.07	\$ 6,071.4
2000	Zucchini		1/2 Bushel (20 lbs.)	1000	\$ 5.99	\$ 5,994.5
						\$ 0.00
						\$ 0.00
						\$ 0.00 \$ 0.00
						\$ 0.00 \$ 0.00 \$ 0.00
						\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00
						\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00
						\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00
					Total Payonus for 1997	\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00
					Total Revenue for 1997	\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 3,465.66
					Total Revenue for 1998	\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 3,465.66
					Total Revenue for 1998 Total Revenue for 1999	\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 3,465.66 \$ 3,465.66
					Total Revenue for 1998	\$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 0.00 \$ 3,465.66 \$ 3,465.66 \$ 6,071.43 \$ 5,994.51 \$ 4,749.31



# Michigan Zucchini WITH Methyl Bromide

Price per Quantity Unit Unit		Total per Acre
Revenue		
Zucchini 1200 1/2 bu \$ 5.99	\$	7,193
Total Revenue	\$	7,193
Expenses		
Field Preparation		
Cover Crop (Materials, Machinery & Labor)	\$	13
Field Tillage (Moldboard plow, Disc twice)	\$	39
Soil Test	\$	2
Lime & Application	\$	13
Fertilizer (Materials)	\$	75
Apply & Incorporate Fertilizer (Machinery & Labor)	\$ \$	• 20
Plastic & Drip Tape (Materials)	\$	284
Lay Plastic & Drip Tape (Machinery & Labor)	\$	98
	\$ \$	341
Fumigate (Materials) Herbicide, Insecticide & Fungicide (Materials)	\$	190
Apply Herbicide, Insecticide & Fungicide (Macha & Lab)	\$	42
•	Ψ	42
Plant & Grow	\$ .	442
Transplants	\$	65
Planting (Machinery & Labor)	\$ \$	53
Cultivate, Hoe & Move Vines (Mach & Lab)	\$	207
Drip Irrigate & Fertigate (Materials, Labor & Electricity)	\$	35
Pollination (Bees)	φ \$	30 30
Scouting & Lab Work		-
Field Maintenance - Driveways & Mowing (Mach & La	\$	7
Harvest	\$	1,440
Pick Crop (Field to Packing Shed)	\$	3,588
Grading & Packing (Includes shipping containers)	Ψ \$	600
Shipping (includes Materials, Machinery & Labor)		647
Sales & Marketing 9% of gross	\$ \$	112
Field Clean-up	\$ \$	75
Management & Supervision	Ф \$	
Interest on Operating Capital 8%	Φ.	144
Land Rent	\$	200
Insurance	\$	8
Other Overhead (Professional Fees, Education & Travel, etc)	\$	10
Total Expenses	\$	8,781
PROFIT	\$	(1,587)

# Michigan Zucchini WITHOUT Methyl Bromide

Price p Quantity Unit U	er nit	Total per Acre
Revenue		
Zucchini 720 1/2 bu \$ 5.4	5 \$	3,920
Total Revenue	\$	3,920
Expenses		
Field Preparation		
Cover Crop (Materials, Machinery & Labor)	\$	13
Field Tillage (Moldboard plow, Disc twice)	\$	39
Soil Test	\$	2
Lime & Application	\$	13
Fertilizer (Materials)	\$	75
Apply & Incorporate Fertilizer (Machinery & Labor)	\$	20
Plastic & Drip Tape (Materials)	\$	284
Lay Plastic & Drip Tape (Machinery & Labor)	\$	98
Fumigate (Materials)	*	•
Herbicide, Insecticide & Fungicide (Materials)	\$	252
Apply Herbicide, Insecticide & Fungicide (Mach & Lab)	\$	54
Plant & Grow	Ψ	01
Transplants	\$	442
Planting (Machinery & Labor)	\$	65
Cultivate, Hoe & Move Vines (Mach & Lab)	\$	53
Drip Irrigate & Fertigate (Materials, Labor & Electricity)		207
Pollination (Bees)	\$	35
Scouting & Lab Work	\$	30
Field Maintenance - Driveways & Mowing (Mach 8		7
Harvest	т СС Ф	,
Pick Crop (Field to Packing Shed)	\$	864
Grading & Packing (Includes shipping containers)	\$	2,153
Shipping (Includes Materials, Machinery & Labor)	\$	360
	φ \$	353
Sales & Marketing 9% of gross Field Clean-up	\$ \$	112
Management & Supervision	φ \$	75
		73 78
Interest on Operating Capital 8% Land Rent	\$ \$	200
	Ψ \$	
Insurance Other Overhead (Professional Fees, Education & Travel, e		8 10
Total Expenses	\$	5,903
PROFIT	\$	(1,982)

Same Strain Strain

For	EPA	Use	Only
			ID#

## Worksheet 3-A(1). Alternatives - Technical Feasibility of Alternatives to Methyl Bromise

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is
not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for
each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: UNEP 1998, B-83, B-281

## Section I. Initial Screening on Technical Feasibility of Alternatives

1.	Are there an	location-specific restrictions that inhibit the use of this alternative on your s	site?
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1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	
,	

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# Worksheet 3-A(1). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

	h Studies on Alternatives to Methyl Bromide
I. Is the study on EPA's website?	
1a. If not on the EPA website, pl	ease attach a copy.
2. Author(s) or researcher(s)	
<del></del>	
3. Publication and Date of Publication	
l. Location of research study	
5. Name of alternative(s) in study. If more	than one alternative, list the ones you wish to discuss.
6. Was crop yield measured in the study?	
7. Describe the effectiveness of the altern	native in controlling pests in the study.
8. Discuss how the results of the study a factors that would affect your adoption	pply to your situation. Would you expect similar results? Are there other of this tool?
	-281 do not indicate that these treatments are effective against P. capsici.
In contrast, many studies indicate that the	e alternatives of chloropicrin and 1,3-Dichloropropene are not effective.

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## Worksheet 3-A(1)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromíde by crop and\_circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional rages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: Evaluation of fungicides for managing Phytophthora crown and fruit rot of zucchini, 2000.

## Section I. Initial Screening on Technical Feasibility of Alternatives

<ol> <li>Are there any location-specific restrictions that inhibit the</li> </ol>	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
****		

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## Worksheet 3-A(1)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright S.D. Linderman Michigan State University Report, 2001 3. Publication and Date of Publication Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35 (1.3-Dichloropropene, chloropicrin) 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. The treatments of Telone C-35 did not offer significant control compared to the untreated control. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? These results are directly applicable to the growing situation in Michigan, USA. It was conducted on the farm of a commercial grower.

#### Worksheet 3-A(1)(b)

ZUCCHINI (Cucurbita Pepo var. Melopepo cv. 'Zucchini' 'Dividend') M.K. Hausbeck, B.D. Cortright, and S.D. Linderman
Phytophthora crown and fruit rot; Phytophthora capsici
Department of Botany and Plant Pathology
Michigan State University
East Lansing, Michigan 48824

EVALUATION OF FUNGICIDES FOR MANAGING PHYTOPHTHORA CROWN AND FRUIT ROT OF ZUCCHINI, 2000: This study was conducted at a cooperator's farm in Cass County, Michigan on a sandy clay loam soil known to have a history of *Phytophthora*, and previously planted to squash. The field was cultivated, bedded, covered with plastic, and drip irrigation installed. Zucchini 'Dividend' was sown on 17 Jun. Plots consisted of one 50-ft row, with 5 ft between rows and 15 in between plants. Weed control, irrigation and fertilization were applied by the grower. Insects were controlled with applications of Provado (3.75 fl oz/A on 23 Jun and 17 Jul), Asana (8 fl oz/A on 27 Jul and 9 Aug), and Endosulfan (32 fl oz/A on 17 Jul). Nova 40WP (5 oz/A) was applied on 27 Jul and 9 Aug for control of powdery mildew. Four treatments were replicated four times in a random rows. Fungicide sprays were applied with a CO<sub>2</sub> backpack boom sprayer equipped with three 11003 nozzles spaced 18 in apart, operating at 60 psi and delivering 50 gal/A. Telone C-35 was applied at bed formation. Acrobat MZ (2.25 lb/A) was applied to replicates 3 and 4 on 25 Jul and 3 Aug. Treatments (except Telone C-35) were applied through the drip irrigation on 5 and 24 Jul. Fruits from the center 5 plants of the treatment row were harvested and weighed three times a week from 27 Jul through 30 Aug, for a total of 15 harvests. Number of infected and total fruit was recorded at harvest and after four days storage at room temperature. Stand count was recorded on 25 Aug.

There were no significant differences among treatments for any parameter measured.

	Stand	Fruit number		_ Total	
Treatment and rate/A, applied at 14-day intervals except fumigant	count 8/25	Infected postharvest (%)	Total <sup>2</sup>	fruit weight (lb)	
Untreated <sup>5</sup>	32.84	11.5	39.0	37.1	
Ultra Flourish 2 pt	36.8	11.6	42.3	38.7	
Telone C-35 at bed formation	34.3	9.9	38.8	36.3	

Stored four days at room temperature; there were significant differences between replicates (Fisher LSD Method; P=0.05). Combined total of 15 harvests; fruit harvested when approximately 12 inches long.

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## Worksheet 3-A(1)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: Evaluation of fumigants for managing Phytophthora crown and fruit rot of summer squash, 1998.

## Section I. Initial Screening on Technical Feasibility of Alternatives

i. Are there	any location-specific restrictions that inhibit the	use of this afternative on your site?	
1a.	Full use permitted	X	
1b.	Township caps	-	
1c.	Alternative not acceptable in consuming country		
1d.	Other (Please describe)		
			<del></del>

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### Worksheet 3-A(1)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright S.D. Linderman 3. Publication and Date of Publication Michigan State University Report, 1999 Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35, Telone C-17 (1,3-Dichloropropene, chloropicrin) 6. Was crop yield measured in the study? % diseased fruit 7. Describe the effectiveness of the alternative in controlling pests in the study. The most effective treatment still had over 50% diseased fruit, which is not commercially acceptable. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? This study is directly applicable, because it was conducted in Michigan, USA on a commercial farm.

#### Worksheet 3-A(1)(c)

SUMMER SQUASH (Curcurbita Pepo 'Seneca Prolific')
Phytophthora Crown and Fruit Rot; Phytophthora capsici

M.K. Hausbeck, B.D. Cortright, and S.D. Linderman Department of Botany and Plant Pathology Michigan State University East Lansing, Michigan 48824

EVALUATION OF FUMIGANTS FOR MANAGING PHYTOPHTHORA CROWN AND FRUIT ROT OF SUMMER SQUASH, 1998: This study was conducted at a cooperator's farm in Cass County, Michigan, on a sandy loam soil known to have disease problems, and previously planted to pumpkin. The field was prepared by plowing, and disking. Fumigant treatments were broadcast on 21 May. The field was bedded, and plugs of summer squash 'Seneca Prolific' were transplanted into holes in the plastic-covered beds. Fertilizer was sidedressed on 24 Jun at 300 lb/A of 6-24-24 and 100 lb/A of 46-0-0. Plots consisted of three 40-ft rows spaced 16 ft apart with a 10-ft wide buffer strip between plots. Treatments were replicated six times in a randomized complete block design. Visual estimations of percentages of diseased plants and diseased fruit were taken on 3 and 12 Aug.

There were no statistical differences in diseased plants among treatments; however, Telone C17 15 gal/A had the least percentage of diseased plants on both observation dates, while Telone C35 10 gal/A had the most percentage of diseased plants. When comparing percentages of diseased fruit, Telone C17 15 gal/A had the least on the first observation date, and by the last observation date of 12 Aug, it had significantly less diseased fruit than any other treatment or the untreated control. However, none of the treatments offered a level of control that was commercially acceptable, since more than half of the fruit were infected.

Treatment and rate	Diseased plants (%) <sup>1</sup>		Diseased fruit (%) <sup>2</sup>	
<del></del>	8/3	8/12	8/3	8/12
Untreated	9.33	90.8	32.0	86.7 b
Telone C17 15 gal/A	5.7	74.5	5.2	53.1 a
Telone C35 10 gal/A	22.8	100.3	32,3	96.0 b
Telone C35 15 gal/A	20.3	82.5	26.7	74.2 b
Telone C35 20 gal/A	17.8	86.6	34.9	80.5 b

Based on a visual estimation of percentage of plant affected.

<sup>&</sup>lt;sup>2</sup>Based on a visual estimation of percentage of fruits infected.

<sup>&</sup>lt;sup>3</sup>Column means with a letter in common or with no letters are not significantly different (Student-Newman-Keuls; P=0.05).

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### Worksheet 3-A(1)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b). When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8. Summarize each of the research studies you cite in the Research Summary Worksheet. If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed. BACKGROUND EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996. There are three major ways you can provide the Agency with proof of your investigative work. (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website. The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area. In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area. Use additional pages as needed. Alternative: 1,3-Dichloropropene, Chloropicrin Study: Evaluation of fumigants for managing Fusarium wilt of muskmelon, 1998. Section I. Initial Screening on Technical Feasibility of Alternatives 1. Are there any location-specific restrictions that inhibit the use of this alternative on your site? 1a. Full use permitted 1b. Township caps 1c. Alternative not acceptable in consuming country 1d. Other (Please describe)

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# Worksheet 3-A(1)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide No X Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright S.D. Linderman 3. Publication and Date of Publication Michigan State University report, 1999 Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35, Telone C-17 (1,3-Dichloropropene, chloropicrin) 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Telone C-17 and C-35 were not effective in limiting crown and root rot. . 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? This study is directly applicable, because it was conducted on a commercial farm in Michigan, USA.

#### Worksheet 3-A(1)(d)

MUSKMELON (Cucumis Melo var. reticulatus 'Quaisar', 'Superstar', 'Rogers ML529Z')

Fusarium Wilt; Fusarium oxysporum f. sp. cucurbitacearum

M.K. Hausbeck, B.D. Cortright, and S.D. Linderman Department of Botany and Plant Pathology Michigan State University East Lansing, Michigan 48824

EVALUATION OF FUMIGANTS FOR MANAGING FUSARIUM WILT OF MUSKMELON, 1998: This study was conducted at a cooperator's farm in Monroe County, Michigan, on a clay loam field known to have disease problems that was previously planted to melons. The field was prepared by plowing and disking. Fumigant treatments (except Methyl Bromide) were broadcast on 18 May. Methyl Bromide was in-row injected, and the field was bedded on 1 Jun. Plugs of three cultivars of muskmelon were transplanted into holes in the plastic-covered beds. Plots consisted of 5-ft rows with 5-ft wide buffer strips between rows. Rows were spaced 6 ft apart with 3.5 ft between plants. Treatments were replicated six times in a randomized complete block design. The field was fertilized to commercial production standards. Stand counts were taken, and plant vigor assessed on 6 Jul. Percent infection was assessed on 11 Aug.

Treatment with methyl bromide completely prevented infection of the muskmelon by *Fusarium oxysporum* f. sp. cucurbitacearum. All other treatments resulted in a minimum of 20% of the plants becoming infected.

Treatment and rate	Infection (%) 8/11	-
Untreated	44.0	
Telone C17 15 gal/A	22.0	
Telone C35 10 gal/A	29.0	
Telone C35 15 gal/A	20.0	. /
Methyl Bromide 6733 200-275 lb/A	0.0	

Based on a scale of 1-5 where 1=poor growth, dead; to 5=green, healthy.

<sup>&</sup>lt;sup>2</sup>Based on visual estimation of percentage of plant affected.

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# Worksheet 3-A(1)(e). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the LPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: Evaluation of fungicides for managing Phytophthora crown and fruit rot of zucchini, 2001.

## Section I. Initial Screening on Technical Feasibility of Alternatives

<ol> <li>Are there any location-specific restrictions that int</li> </ol>	hibit the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	-
1c. Alternative not acceptable in consuming co	ountry
1d. Other (Please describe)	

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## Worksheet 3-A(1)(e). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright 3. Publication and Date of Publication Michigan State University report, 2002 4. Location of research study Michigan, USA 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35 (1,3-Dichloropropene, chloropicrin) 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Telone C-35 did not provide a benefit compared to the untreated control. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? This study is directly applicable to the current situation in Michigan, USA, since it was conducted with a commercial grower.

#### Worksheet 3-A(1)(e)

ZUCCHINI (Cucurbita Pepo var. Melopepo cv. 'Zucchini' 'Spineless Beauty')
Phytophthora crown and fruit rot; Phytophthora capsici

M.K. Hausbeck, and B.D. Cortright Department of Plant Pathology Michigan State University East Lansing, Michigan 48824

#### Evaluation of fungicides for managing Phytophthora crown and fruit rot of zucchini, 2001.

This study was conducted at a cooperator's farm in Cass County, Michigan on a sandy clay loam soil known to have a history of *Phytophthora*, and previously planted to squash. The field was cultivated, bedded, covered with plastic, and drip irrigation installed. Zucchini 'Spineless Beauty' was sown on 30 May. Plots consisted of one 20-ft row, with 5 ft between rows and 18 in between plants. Weed control, irrigation and fertilization were applied by the grower. Insects were controlled with applications of Pounce (8.0 fl oz/A) and Provado (3.75 fl oz/A) on 9 Jul. Five treatments were replicated four times in a random rows. Treatments (except Telone C-35) were applied through the drip irrigation on 19 Jun and 16 Jul. Fruits from the center 5 plants of the treatment row were harvested and weighed twelve times during the period from 3 Jul through 30 Jul. Number of infected and total fruit was recorded at harvest and after four days storage at room temperature. Stand count was recorded on 27 Jul.

There were no significant differences among treatments for any parameter measured.

Treatment and rate/A, applied at 28-day intervals except fumigant	Stand count 7/27	Number of healthy fruit post harvest	Total fruit weight (lb)
Untreated	14.0	27.5	26.1
Ultra Flourish 2 pt	13.5	17.3	19.3
Deny 2 pt	14.0	15.0	17.6
Root Guard Plus 17 oz	14.0	22.5	26.1
Telone C-35 at bed formation 25 gal	14.0	11.0	9.8

Stored four days at room temperature; there were significant differences between replicates (Student Newman Keuls; P=0.05).

<sup>&</sup>lt;sup>2</sup>Combined total of 12 harvests; fruit harvested when approximately 12 inches long.

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### Worksheet 3-A(1)(f). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: Alternatives for methyl bromide on watermelon, 2002.

## Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions	that inhibit the use of this alternative on your site?
1a. Full use permitted	X

1b.	Township caps	
1c.	Alternative not acceptable in consuming country	

1d.	Other (Please describe)	

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## Worksheet 3-A(1)(f). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide No X 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) M.K. Hausbeck E. Kerlikowski B.D. Cortright 3. Publication and Date of Publication Research in progress 4. Location of research study Michigan, USA 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35 (1,3-Dichloropropene, chloropicrin) 6. Was crop yield measured in the study? Yes \_\_\_\_ 7. Describe the effectiveness of the alternative in controlling pests in the study. The plants in the Telone C-35 treatment were stunted, with a reduction in leaf size and canopy, resulting in sunburn on fruit. The plants in the methyl bromide treatment were full-sized and grew vigorously. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable, since the research was conducted in Michigan, USA.

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### Worksheet 3-A(1)(g). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area:) The Agency has developed-a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-Dichloropropene, Chloropicrin Study: Alternatives for methyl bromide on cucurbits, 2002.

## Section I. Initial Screening on Technical Feasibility of Alternatives

i. Are there any location-specific restrictions that inhibit the i	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
		<del></del>

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# Worksheet 3-A(1)(g). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

3e	ection II. Existing Res	search S	Studies on Alf	ternative	es to Methyl	Bromide
1.	Is the study on EPA's website?	•	Yes	No_	X	
	1a. If not on the EPA we	bsite, pleas	se attach a copy.			
2.	Author(s) or researcher(s)	M.K. Haus	beck			
		B.D. Cortri	ght			
3.	Publication and Date of Publica	ation	Research in progres	ss		
4.	Location of research study	Michigan,	USA			
5	Name of alternative(s) in study	If more tha	an one alternative. li	st the ones	you wish to discus	<b>5</b> S.
٥.	Telone C-35, Chloropicrin 100%,					
					-	
6.	Was crop yield measured in the	study?	Yes	No_	X	
7.	Describe the effectiveness of the		ve in controlling pes	sts in the stu	ıdy.	
	Fields have not been harvested y	et.				
					1-	
_	Discuss how the results of the					
8.	other factors that would affect	study appi your adopt	ion of this tool?	vvoulu you t	expect similar rest	iits: Are there
	The results of this study are direct	tly applicab	le, since the research	was conduc	ted in Michigan, US	5A.
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### Worksheet 3-A(2). Alternatives - Technical Feasibility of Alternatives to Methyl Booking

In this worksheet, you should address why an alternative pest management strategy on the list (see previous not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of work each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: 1,3-D, Metam Sodium, Basamid	Study: UNEP 1998, B-281

## Section I. Initial Screening on Technical Feasibility of Alternatives

<ol> <li>Are there any location-specific restrictions that inhibit the</li> </ol>	e use of this alternative on your site?	
1a. Fuli use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		

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# Worksheet 3-A(2). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

<b>Se</b>	ection II. Existing Research Studies on Alternatives to Methyl Bromide
1.	Is the study on EPA's website?  Yes X  No No
	1a. If not on the EPA website, please attach a copy.
2.	Author(s) or researcher(s)
3.	Publication and Date of Publication
4.	Location of research study
5.	Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss.  1,3-D, Metam Sodium, Basamid
	1,5-D, Welah Sodian, Dasamo
6.	Was crop yield measured in the study? Yes No
_	The state of the s
7.	Describe the effectiveness of the alternative in controlling pests in the study.
	Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool?
	The study in UNEP 1998 B-281 indicates that direct injection of metam sodium in bands to soil does not provide
	consistent control due to non-uniform distribution in the soil. Also, problems with microorganisms that degrade the
	chemical, thereby making it less effective, have been noted.

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# Worksheet 3-A(2)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.
For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).
When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the Research Summary Worksheet.
If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.
BACKGROUND
EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996
There are three major ways you can provide the Agency with proof of your investigative work.
<ul><li>(1) Conduct and submit your own research</li><li>(2) Cite research that has been conducted by others</li><li>(3) Cite research listed on the EPA website</li></ul>
Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.
The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.
In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.
Use additional pages as needed.
Alternative: 1,3-D, Metam Sodium, Basamid Study: Alternatives for methyl bromide on cucurbits and
solanaceous crops, 2002.
Section I. Initial Screening on Technical Feasibility of Alternatives
1. Are there any location-specific restrictions that inhibit the use of this alternative on your site?
1a. Full use permitted X
1b. Township caps
1c. Alternative not acceptable in consuming country
1d. Other (Please describe)

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# Worksheet 3-A(2)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes No X 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright 3. Publication and Date of Publication Research in progress 4. Location of research study Michigan, USA 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Multigard FFA, Multigard Protect, Multigard Protect + Vapam HL, CX-100 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Fields have not been harvested yet. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable, since the research was conducted in Michigan, USA.

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Worksheet 3-A(2)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide
In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.
For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).
When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the Research Summary Worksheet.
If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.
BACKGROUND
EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.
There are three major ways you can provide the Agency with proof of your investigative work.  (1) Conduct and submit your own research  (2) Cite research that has been conducted by others  (3) Cite research listed on the EPA website
Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.
The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.
In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.
Use additional pages as needed.
Alternative: 1,3-D, Metam Sodium Study: Alternatives for methyl bromide on cucurbits, 2002.
Section I. Initial Screening on Technical Feasibility of Alternatives
1. Are there any location-specific restrictions that inhibit the use of this alternative on your site?
1a. Full use permittedX
1b. Township caps
1c. Alternative not acceptable in consuming country
1d. Other (Please describe)

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# Worksheet 3-A(2)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

ection II. Existing Res	search	Studies on A	Alternatives	to Methyl Bı	romide
1. Is the study on EPA's website	?	Yes	No	<u> </u>	
1a. If not on the EPA we	bsite, ple	ase attach a copy.			<i>2</i> 1
2. Author(s) or researcher(s)	M.K. Hau	usbeck			
	B.D. Cort	tright		- 44-	
3. Publication and Date of Public	ation	Research in prog	ress		
I. Location of research study	Michigan	, USA			
5. Name of alternative(s) in study Telone C-35, Chloropicrin 100%,	lodometha		cken manure		
. Was crop yield measured in th	e study?	Yes	No	<u> </u>	
. Describe the effectiveness of t Fields have not been harvested y		tive in controlling p			
. Discuss how the results of the other factors that would affect	your adop	otion of this tool?	ո. Would you exp	ect similar results	
The results of this study are direct	tly applica	ble, since the resear	rch was conducted	in Michigan, USA.	***
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### Worksheet 3-A(3). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8,

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Metam Sodium, Crop Rotation	Study: UNEP 1998, B-39, B-74	
, titoritativo: inotati o o o o o o o o o o o o o o o o o o o	July: 1000, B 00, B 14	•

### Section I. Initial Screening on Technical Feasibility of Alternatives

. Are there any location-specific restrictions that inhibit the	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		

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# Worksheet 3-A(3). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes \_\_X \_\_ No \_\_\_\_\_ 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) \_\_\_\_\_\_ 3. Publication and Date of Publication \_\_\_\_\_\_ 4. Location of research study \_\_\_\_\_\_ 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. \_\_\_\_\_\_\_ Metam Sodium, Crop Rotation \_\_\_\_\_\_\_ 6. Was crop yield measured in the study? Yes \_\_\_\_\_\_\_ No \_\_\_\_\_\_\_ 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The efficacy of crop rotation depends on the life cycle of the pathogen and its ability to overwinter and persist in soils. Phytophithora capsici has an overwintering structure called an oospore that is capable of surviving for long periods of time, thereby negating the benefits of crop rotation. Metam sodium is not considered a good product

against fungi, especially *Phytophthora*, but rather is used for weed control and for nematode problems. The rotation suggested in this study primarily lists crops that are susceptible to *P. capsici*, including pepper, cucumber,

tomato and squash. Using this suggested rotation would exacerbate the disease problem. Therefore, this

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alternative would not be effective for Michigan growers.

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			ID#		

### Worksheet 3-A(4). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete question

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible—alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Biofumigation	Study: UNEP 1998, B-83, B-41, B-91, B-94
3	

### Section I. Initial Screening on Technical Feasibility of Alternatives

. Are there any location-specific restrictions that inh	ibit the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming con	untry
. 1d. Other (Please describe)	

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# Worksheet 3-A(4). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes X No \_\_\_\_ 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Biofumigation 6. Was crop yield measured in the study? Yes 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Biofumigation does not readily apply to Michigan's situation. The reasons for this include the lack of evidence that this treatment works for Phytophthora capsici under Michigan's cool climate. Phytophthora's oospore would not be

killed using biofumigation in Michigan's soils. Beneficial predatory nematodes important for biofumigation have not

been identified or quanitified in Michigan's vegetable growing regions.

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worksneet 3-A(4)(b). Alternatives - Technica	i reasibility of Alternatives to Methyl Bromide
In this worksheet, you should address why an alternative pe not effective for your conditions. This worksheet contains 9 each research study you use to evaluate a single methyl bro	st management strategy on the list (see previous page) is or is questions. You must complete one copy of worksheet 3-A for mide alternative. Use additional pages as need.
the worksheet 3-A(1)(c). For the second alternative, first resident alternative, second research study, label the worksheet 3-(A)	, first research study, label the worksheet 3-A(1)(a). For the et 3-A(1)(b). For the first alternative, third research study, label earch study, label the worksheet 3-(A)(2)(a). For the second ((2)(b).
When completing Section II, if you cite a study that is on the	EPA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the Rese	-
If you prefer, you may provide the information requested in t research reports. The narrative review must reply to Section Worksheet of relevant treatments should be provided for each	I and questions 1 through 8 in Section II. A Research Summary
BACKGROUND	
EPA must consider whether alternative pest control measures (pesuccessfully instead of methyl bromide by crop and circumstance alternative pest control regimens for various crops, which can be	esticide and non-pesticidal, and their combination) could be used (geographic area.) The Agency has developed a list of possible found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.
There are three major ways you can provide the Agency with produced the	of of your investigative work.
<ul><li>(1) Conduct and submit your own research</li><li>(2) Cite research that has been conducted by others</li><li>(3) Cite research listed on the EPA website</li></ul>	·
Whether you conduct the research yourself or cite studies develop scientifically sound manner. The studies should include a descript application intervals, pest pressure, weather conditions, varieties outcome. You must submit copies of each study to EPA unless	tion of the experimental methodology used, such as application rates, of the crop used, etc. All results should be included, regardless of
The Agency has posted many research studies on a variety of cro EPA will add studies to its website as they become publicly availa websites for studies that pertain to your crop and geographic area	ps on its website and knows of more studies currently in progress. ble. You are encouraged to review the EPA website and other .
the Agency and explain why they cannot be used for your crop an	e in Seattle). You should look at the list of alternatives provided by d in your geographic area.
Use additional p	pages as needed.
	y: Alternatives for methyl bromide on cucurbits and solanaceous crops, 2002.
ection I. Initial Screening on Technical F	easibility of Alternatives
4. And the recognition of the second	
<ol> <li>Are there any location-specific restrictions that inhibit the unit is a full use permitted</li> </ol>	
1b. Township caps	X
1c. Alternative not acceptable in consuming country	
Alternative not acceptable in consuming country     d. Other (Please describe)	
id. Other (Flease describe)	,

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# Worksheet 3-A(4)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes \_\_\_\_\_ No \_\_X\_\_\_

1a. If not on the EPA website, please attach a copy.				
2.	Author(s) or researcher(s)	M.K. Hausbeck		
		B.D. Cortright		
3.	Publication and Date of Publication	ation Research in progress		
4.	Location of research study	Michigan, USA		
5.	5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss.  Multigard FFA, Multigard Protect, Multigard Protect + Vapam HL, CX-100			
6.	Was crop yield measured in the	e study? Yes No X		
7. Describe the effectiveness of the alternative in cont		he alternative in controlling pests in the study.		
	Fields have not been harvested yet.			
	<u> </u>			
8.	other factors that would affect			
	The results of this study are direct	ctly applicable, since the research was conducted in Michigan, USA.		
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### Worksheet 3-A(4)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b). When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8. Summarize each of the research studies you cite in the Research Summary Worksheet. If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed. BACKGROUND EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996. There are three major ways you can provide the Agency with proof of your investigative work. (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website. The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area. In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area. Use additional pages as needed. Alternative: Biofumigation Study: Alternatives for methyl bromide on cucurbits, 2002. Section I. Initial Screening on Technical Feasibility of Alternatives 1. Are there any location-specific restrictions that inhibit the use of this alternative on your site? 1a. Full use permitted 1b. Township caps 1c. Alternative not acceptable in consuming country 1d. Other (Please describe)

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# Worksheet 3-A(4)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to: Methyl Bromide 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright 3. Publication and Date of Publication Research in progress Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35, Chloropicrin 100%, lodomethane, Composted chicken manure 6. Was crop yield measured in the study? No X 7. Describe the effectiveness of the alternative in controlling pests in the study. Fields have not been harvested yet. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable, since the research was conducted in Michigan, USA.

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# Worksheet 3-A(5). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Solarization	Study:	<b>UNEP 19</b>	5. UNEP	1998	B-83	B-281	Δ-77	R-91	R-94
Alternative: Goldinzation	otuay.	OHILL 13	Jo, OILL	1330	,,	, 0-201,	Α-ιι,	D-31,	D-34

# Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the use of this alternative on your site?

<ul> <li>Alternative not acceptable in consuming count</li> </ul>	у
d. Other (Please describe)	

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# Worksheet 3-A(5). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

Section II.	Existing Research Studies on Alternatives to Methyl Bromide
1. Is the study of	n EPA's website? Yes X No No
1a. If r	ot on the EPA website, please attach a copy.
2. Author(s) or i	researcher(s)
3. Publication a	nd Date of Publication
4. Location of re	esearch study
5. Name of alter Solarization	native(s) in study. If more than one alternative, list the ones you wish to discuss.
6. Was crop yiel	d measured in the study? Yes No
7. Describe the	effectiveness of the alternative in controlling pests in the study.
	the results of the study apply to your situation. Would you expect similar results? Are there That would affect your adoption of this tool?
	the soil to kill the overwintering spores is not feasible in a northern state where the growing season
is short (May to	September), and cold temperatures (<50 F) prevail through much of the year. Also, this

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# Worksheet 3-A(6). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) courselessfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Steam	Study: UNEP 1998, B-83, B-90, B-91, B-282	
Section I. Initial Screening	on Technical Feasibility of Alternatives	
1. Are there any location-specific restri	ictions that inhibit the use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in	consuming country	
1d. Other (Please describe)		

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# Worksheet 3-A(6). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

Se	ction II. Existing Research Studies on Alternatives to Methyl Bromide
1.	Is the study on EPA's website?
	1a. If not on the EPA website, please attach a copy.
2.	Author(s) or researcher(s)
3.	Publication and Date of Publication
4.	Location of research study
5.	Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss.  Steam
6.	Was crop yield measured in the study? Yes No
7.	Describe the effectiveness of the alternative in controlling pests in the study.
8.	Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool?
	In the studies referenced, steaming has been used in protected production systems, such as greenhouses. The
	use of steam has not proven economical and practical when large, unprotected areas are treated. In Michigan
	systems, Phytophthora capsici has an airborne spore that would render the use of steam ineffective.

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### Worksheet 3-A(7). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative:	Biological Control	Study: UNEP 1998, B-83, B-45, B-91, B-92
Section I.	Initial Screening on Technic	cal Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the	use of this alternative on	your site?
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
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# Worksheet 3-A(7). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

CHOIL III EXIONIS LICEUM S	h Studies on Alternatives to Methyl Bromide
Is the study on EPA's website?	YesX No
1a. If not on the EPA website, p	olease attach a copy.
Author(s) or researcher(s)	
Publication and Date of Publication	
Location of research study	
	re than one alternative, list the ones you wish to discuss.
	<u> </u>
Was crop yield measured in the study	/? Yes No
	rnative in controlling pests in the study.
Discuss how the results of the study a other factors that would affect your ad	apply to your situation. Would you expect similar results? Are there doption of this tool?
	capsici was not a target pathogen, so they do not apply to Michigan's

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### Worksheet 3-A(7)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Section I. Initial Screening on Technical Feasibility of Alternatives  1. Are there any location-specific restrictions that inhibit the use of this alternative on your site?  1a. Full use permitted  2x  1b. Township caps  1c. Alternative not acceptable in consuming country  1d. Other (Please describe)	Alternative: Biological Control	Study: Evaluation of fungicides for managing Phytophtho crown and fruit rot of zucchini, 2001.	ra
1a. Full use permitted X  1b. Township caps  1c. Alternative not acceptable in consuming country	Section I. Initial Screening on I		
1b. Township caps  1c. Alternative not acceptable in consuming country	1. Are there any location-specific restrictions	that inhibit the use of this alternative on your site?	
1c. Alternative not acceptable in consuming country	1a. Full use permitted	X	
·	1b. Township caps		
1d. Other (Please describe)	1c. Alternative not acceptable in consu	uming country	
	1d. Other (Please describe)		

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# Worksheet 3-A(7)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? No X Yes 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) M.K. Hausbeck B.D. Cortright 3. Publication and Date of Publication Michigan State University report, 2001 4. Location of research study Michigan 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Biological Control Pseudomonas bacterium, Trichoderma harzianum fungus 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. The treatments included in the study were not significantly different from the untreated control. The biocontrol agents of Pseudomonas and Trichoderma did not offer a benefit compared to the untreated control. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Since this study was conducted in Michigan, the results are directly applicable. Biocontrol agents at this time do not offer a viable control alternative for Michigan cucurbit growers.

### Worksheet 3-A(7)(b)

ZUCCHINI (Cucurbita Pepo var. Melopepo cv. 'Zucchini' 'Spineless Beauty')
Phytophthora crown and fruit rot; Phytophthora capsici

M.K. Hausbeck, and B.D. Cortright Department of Plant Pathology Michigan State University East Lansing, Michigan 48824

### Evaluation of fungicides for managing Phytophthora crown and fruit rot of zucchini, 2001.

This study was conducted at a cooperator's farm in Cass County, Michigan on a sandy clay loam soil known to have a history of *Phytophthora*, and previously planted to squash. The field was cultivated, bedded, covered with plastic, and drip irrigation installed. Zucchini 'Spineless Beauty' was sown on 30 May. Plots consisted of one 20-ft row, with 5 ft between rows and 18 in between plants. Weed control, irrigation and fertilization were applied by the grower. Insects were controlled with applications of Pounce (8.0 ft oz/A) and Provado (3.75 ft oz/A) on 9 Jul. Five treatments were replicated four times in a random rows. Treatments (except Telone C-35) were applied through the drip irrigation on 19 Jun and 16 Jul. Fruits from the center 5 plants of the treatment row were harvested and weighed twelve times during the period from 3 Jul through 30 Jul. Number of infected and total fruit was recorded at harvest and after four days storage at room temperature. Stand count was recorded on 27 Jul.

There were no significant differences among treatments for any parameter measured.

Treatment and rate/A, applied at 28-day intervals except fumigant	Stand count 7/27	Number of healthy fruit post harvest	Total fruit weight (lb)
Untreated	14.0	27.5	26.1
Ultra Flourish 2 pt	13.5	17.3	19.3
Deny 2 pt	14.0	15.0	17.6
Root Guard Plus 17 oz	14.0	22.5	26.1
Telone C-35 at bed formation 25 gal	14.0	11.0	9.8

Stored four days at room temperature; there were significant differences between replicates (Student Newman Keuls; P=0.05).

<sup>&</sup>lt;sup>2</sup>Combined total of 12 harvests; fruit harvested when approximately 12 inches long.

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# Worksheet 3-A(7)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

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Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Biological Control	Study: Evaluation of materials for control of Phytophthora crown rot of summer squash, 1998.
Section I. Initial Screening on Technic	al Feasibility of Alternatives
Are there any location-specific restrictions that inhib	it the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming coun	try
1d. Other (Please describe)	

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# Worksheet 3-A(7)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

1. Is the study on EPA's website?	Yes NoX
1a. If not on the EPA we	bsite, please attach a copy.
2. Author(s) or researcher(s)	G.J. Holmes
	M.E. Lancaster
	D.W. Pollard
3. Publication and Date of Publica	Fungicide and Nematicide Tests 54:240, 1999
4. Location of research study	Hendersonville, North Carolina
	. If more than one alternative, list the ones you wish to discuss.
Biological Control	
Clicaladium virana Durkhaldaria	connaio
Gliocladium virens, Burkholderia	cepacia
6. Was crop yield measured in the	e study?  Yes X No  he alternative in controlling pests in the study.
6. Was crop yield measured in the	e study? Yes X No
6. Was crop yield measured in the 7. Describe the effectiveness of the The biocontrol treatment was con	e study?  Yes X No No he alternative in controlling pests in the study.
6. Was crop yield measured in the 7. Describe the effectiveness of the The biocontrol treatment was con	e study? Yes X No he alternative in controlling pests in the study.
6. Was crop yield measured in the 7. Describe the effectiveness of the The biocontrol treatment was condisease suppression.	the alternative in controlling pests in the study.  Inparable to the untreated control and did not provide a significant benefit in  Study apply to your situation. Would you expect similar results? Are there other
6. Was crop yield measured in the 7. Describe the effectiveness of the The biocontrol treatment was condisease suppression.  8. Discuss how the results of the factors that would affect your a	the alternative in controlling pests in the study.  Inparable to the untreated control and did not provide a significant benefit in  Study apply to your situation. Would you expect similar results? Are there other

G. J. Holmes<sup>1</sup>, M. E. Lancaster<sup>2</sup>, and D. W. Pollard<sup>2</sup>, <sup>1</sup>Dept Plant Pathology, NC State Univ, Box 7616, Raleigh, NC 27695, and <sup>2</sup>NC Coop. Ext. Service, Hendersonville, NC 28792

EVALUATION OF MATERIALS FOR CONTROL OF PHYTOPHTHORA CROWN ROT OF SUMMER

SQUASH, 1998: The experiment was conducted in a commercial squash field near Hendersonville NC (GPS coordinates: N35°18.047; W082°23.872) with a known history of Phytophthora crown rot (PCR). Field soil type was a Delanco B loam (pH 6.6). Applications of lime (1000 lb/Ac) and fertilizer (500 lb/Ac of 10-20-20) were made on 09 May. Seed was planted on 22 May into flat soil and thinned to 24-in spacing on 16 Jun. All seed was treated with Captan except that which was treated with the bacterium (Burkholderia cepacia) and the fungus (Gliocladium virens) and a nontreated control. In both treatments, Captan was removed from the seed by washing in water for 10 min. The B. cepacia + G. virens seed treatment was applied to seed at the Biocontrol of Plant Diseases Laboratory, USDA, Beltsville, MD. All seed used was from the same seed lot. Fertilizer was sidedressed (230 lb of 15-0-14) on 13 Jun. No irrigation was used. Treatments were randomized in 4 complete blocks. Plots were single rows on 36-in. spacing and 35 ft long with 5-ft fallow borders on each end and separated by a non-treated row on each side. Fungicide treatments were applied using a manual, diaphram-type Solo backpack sprayer equipped with a single nozzle (Conejet TXVS-26) and operating at 40 psi (45 gal/A). To ensure adequate coverage, treatments were applied by making a pass on each side of the treated row (2 passes per plot). Treatments were applied on a 7-day interval with applications on 11, 18, 25 Jun, 01, 09, and 15 Jul. Disease incidence (% mortality due to PCR) was counted weekly (11, 18, 25 Jun, 01, 09, 15 and 23 Jul). Fruit were picked weekly (01, 09, 15, and 23 Jul).

PCR was detected in an adjacent field of squash prior to emergence in test plots. The original stand was excellent, but plants began dying shortly after the first treatment applications (4 true leaves). While other pathogens may cause damping-off in squash (e.g., Rhizoctonia and Pythium), the only pathogen detected in periodic spot checks was P. capsici. Disease distribution was both clustered and scattered. Disease pressure was considered very high and conditions were conducive for disease development, especially during the first half of the experiment. Due to clusters of high disease incidence, variability between replicate treatments was high, yielding very little statistical separation of treatments, but significant block effects. No treatment provided a commercially acceptable level of control. The Ridomil Gold Bravo treatment yielded the lowest percent mortality at each evaluation, but final mortality was 61% in these plots. Differences in yield were not significant.

·			%	Mortality	7			
Treatment and rate/A	11 Jun	18 Jun	25 Jun	1 Jul	9 Jul	15 Jul	23 Jul	AUDPC 1
Ridomil Gold Bravo, 2 lb	0.0	2.38	13.0	27.8	31.3	50.1	61.3	11.1 a <sup>2</sup>
Dithane 75DF, 3.2 lb	0.0	8.7	19.0	36.2	47.0	53.0	71.0	14.2 ab
Acrobat MZ, 2.25 lb (alternate stem base-foliar) <sup>3</sup>	2.3	13.9	19.7	32.9	40.0	54.9	69.5	14.3 ab
Acrobat MZ, 2.25 lb	- 1.1	5.7	29.6	53.5	62.7	66.1	70.8	18.1 ab
Acrobat 50WP, 3.22 lb + Kocide 2000, 1.86 lb	1.1	20.7	37.3	52.3	60.6	64.3	82.0	19.7 ab
Tattoo C, 1.47 lb	0.0	0.0	23.4	55.1	77.0	88.5	89.8	20.6 ab
Aliette WDG, 5 lb	0.0	3.5	17.4	60.3	92.2	95.9	100	22.8 ab
Quadris, 0.87 lb	3.4	26.6	42.9	63.8	75.0	82.2	88.7	24.1 ab
Nontreated (seed w/ Captan)	4.8	29.8	51.3	65.6	79.0	88.2	92.1	26.0 ab
Nontreated (seed w/o Captan)	6.1	28.0	36.7	76.7	91.7	95.8	98.6	27.6 b
Seed treatment (Gliocladium virens + Burkholderia cepacia)	3.1	37.2	54.4	81.0	89.3	98.2	100	29.4 b
LSD (P=0.05)	ns	ns	ns	ns	43.6	39.6	30.0	15.3

AUDPC = Area Under the Disease Progress Curve.

F&N Tests 54:240

<sup>&</sup>lt;sup>2</sup> Values are the means of 4 replicate plots; values followed by the same letter within a column are not significantly different (K=100, Duncan-Waller K-ratio test).

<sup>&</sup>lt;sup>3</sup> application alternated weekly between a stem-base directed spray and a standard foliar spray.

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### Worksheet 3-A(7)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

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- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Biological Control	Study: Alternatives for methyl bromide on cucurbits, 2002.
Section I. Initial Screening on Technic	cal Feasibility of Alternatives
Are there any location-specific restrictions that inhib	it the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming coun	
1d. Other (Please describe)	

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# Worksheet 3-A(7)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

I. Is the study on EPA's website?	Yes	No	X	
1a. If not on the EPA we	bsite, please attach a copy.			
Author(s) or researcher(s)	M.K. Hausbeck			
	B.D. Cortright			
Publication and Date of Publication	Research in prog	ess		
Location of research study	Michigan, USA			
Telone C-35, Chloropicrin 100%,	lodomethane, Composted chic	ken manure	· · · · · · · · · · · · · · · · · · ·	
Telone C-35, Chloropicrin 100%,  Was crop yield measured in the  Describe the effectiveness of the	e study?  Yes  The alternative in controlling p	ken manure	X	
Telone C-35, Chloropicrin 100%,  Was crop yield measured in the  Describe the effectiveness of the Fields have not been harvested y	e study?  Yes  The alternative in controlling points.	Noests in the stud	X dy.	
Was crop yield measured in the Describe the effectiveness of the Fields have not been harvested y	e study?  Yes  The alternative in controlling post.  Study apply to your situation your adoption of this tool?	Noests in the stud	X dy.	ar results? Are ther

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### Worksheet 3-A(8). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b). When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8. Summarize each of the research studies you cite in the Research Summary Worksheet. If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed. **BACKGROUND** EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used. successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996. There are three major ways you can provide the Agency with proof of your investigative work. (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website. The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area. In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area. Use additional pages as needed. Alternative: Cover Crops, Mulching Study: UNEP 1998, B-91 Section I. Initial Screening on Technical Feasibility of Alternatives 1. Are there any location-specific restrictions that inhibit the use of this alternative on your site? 1a. Full use permitted 1b. Township caps 1c. Alternative not acceptable in consuming country 1d. Other (Please describe)

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# Worksheet 3-A(8). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

Section II. Existing Research	Studies on Alternatives to Methyl E	Bromide
1. Is the study on EPA's website?	Yes X No	
1a. If not on the EPA website, ple	ease attach a copy.	
2. Author(s) or researcher(s)		
3. Publication and Date of Publication		
4. Location of research study		
	han one alternative, list the ones you wish to discuss	
Carran Carran Adulation		-
6. Was crop yield measured in the study?	Yes No	
7. Describe the effectiveness of the alternation		/
	·	
8. Discuss how the results of the study apportant other factors that would affect your adoption.	oly to your situation. Would you expect similar resultention of this tool?	s? Are there
The results of the study are not relative to the	ne situation in Michigan, because the examples provided	specifically
	The only pathogen included was Sclerotinia sclerotiorum.	
growers are managing Phytophthora capsic	i currently using black plastic mulch, but it is not a viable	alternative
alone to control this pathogen.		

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### Worksheet 3-A(8)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the

worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

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- (3) Cite research listed on the EPA website

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The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Cover Crops, Mulching	Study:	Watermelons in Mexico:	Mulch	and Soil	<b>Amendments</b>

### Section I. Initial Screening on Technical Feasibility of Alternatives

1. A	Are there any location-specific restrictions that inhibit	the use of this alternative on your site	?
	1a. Full use permitted	X	
	1b. Township caps		
	1c. Alternative not acceptable in consuming countr	y	
	1d. Other (Please describe)		
			***************************************

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# Worksheet 3-A(8)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

. Is the study on EPA's website?	Yes X No
1a. If not on the EPA website, p	please attach a copy.
. Author(s) or researcher(s)	
. Publication and Date of Publication	
. Location of research study	
	re than one alternative, list the ones you wish to discuss.
. Was crop yield measured in the study	y? Yes No
7. Describe the effectiveness of the alter	ernative in controlling pests in the study.
	apply to your situation. Would you expect similar results? Are there ot
B. Discuss how the results of the study a factors that would affect your adoption	apply to your situation. Would you expect similar results? Are there ot
8. Discuss how the results of the study of factors that would affect your adoption.  The study in Mexico does not apply to the	apply to your situation. Would you expect similar results? Are there ot on of this tool?

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### Worksheet 3-A(8)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b). When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8. Summarize each of the research studies you cite in the Research Summary Worksheet. If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed. BACKGROUND EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996. There are three major ways you can provide the Agency with proof of your investigative work. (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates. application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website. The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area. In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area. Use additional pages as needed. Alternative: Cover Crops, Mulching Study: Alternatives for methyl bromide on cucurbits and solanaceous crops, 2002. Section I. Initial Screening on Technical Feasibility of Alternatives 1. Are there any location-specific restrictions that inhibit the use of this alternative on your site? 1a. Full use permitted 1b. Township caps 1c. Alternative not acceptable in consuming country 1d. Other (Please describe)

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# Worksheet 3-A(8)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

Se	ection II. Existing Res	earch St	tudies on A	lternative	es to Methy	I Bromide
1.	is the study on EPA's website?	•	Yes	No_	<u> </u>	
	1a. If not on the EPA we	bsite, please	attach a copy.			
2.	Author(s) or researcher(s)	M.K. Hausbe	eck			
		B.D. Cortright	nt			
3.	Publication and Date of Publica	ation <u></u>	Research in prog	ress		
4.	Location of research study	Michigan, US	SA		\\	
	Name of alternative(s) in study.					
Э.	Multigard FFA, Multigard Protect,	Multigard Pro	tect + Vapam HL	., CX-100	, ou 111511 to alou	
6.	Was crop yield measured in the	study?	Yes	No_	X	
7.	Describe the effectiveness of the Fields have not been harvested y	-4				
	Fleids flave not been flatvested y	<u> </u>				
8.	Discuss how the results of the other factors that would affect			. Would you	expect similar re	sults? Are there
	The results of this study are direct	tly applicable,	since the resear	ch was conduc	ted in Michigan, L	JSA.
	1845 1846 1846					

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### Worksheet 3-A(8)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combi<u>nation</u>) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

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- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Cover Crops, Mulching	Study: Alternatives for methyl bromide on cucurbits, 2002.

### Section I. Initial Screening on Technical Feasibility of Alternatives

1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	<u></u>
1d. Other (Please describe)	·

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# Worksheet 3-A(8)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

1. Is the study on EPA's website	? Yes	No _	X	J
1a. If not on the EPA w	ebsite, please attach a copy	<i>1</i> .		
2. Author(s) or researcher(s)	M.K. Hausbeck			
	B.D. Cortright			
. Publication and Date of Public	Research in pro	ogress		
. Location of research study	Michigan, USA			
		hickon manusa		
Telone C-35, Chloropicrin 100%	, lodomethane, Composted cl	nicken manure		-
			· x	
. Was crop yield measured in th	e study? Yes he alternative in controlling	No		
. Was crop yield measured in th	e study? Yes he alternative in controlling	No pests in the stu	dy.	
. Was crop yield measured in the Describe the effectiveness of the Fields have not been harvested.	e study? Yes he alternative in controlling yet. study apply to your situatio	No_ pests in the stu	dy.	
. Was crop yield measured in the Describe the effectiveness of the Fields have not been harvested.	e study? Yes he alternative in controlling yet.  study apply to your situation your adoption of this tool?	No pests in the stur / on. Would you e	dy.	results? Are there

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### Worksheet 3-A(9). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, reconditions. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Crop Rotation, Fallow . Study:	UNEP 1998, B-94, B-91, B-83, B-282
---	------------------------------------

### Section I. Initial Screening on Technical Feasibility of Alternatives

Are there any location-specific restrictions that inhibit the	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		

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		ID#	

## Worksheet 3-A(9). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? No 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Crop Rotation, Fallow 6. Was crop yield measured in the study? Yes 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Crop rotation to crops not susceptible to crown, root and fruit rot caused by Phytophthora capsici is practiced routinely by growers of cucurbit crops in Michigan. This management practice is not adequate, because of the long-lived oospore of this pathogen. Since many other vegetable crops are also susceptible, including all Solanaceous crops and beans (new report of lima beans as a host), this would make rotation difficult even if it was effective. Crop rotation and fallow is not a suitable alternative to manage P. capsici on cucurbits in Michigan.

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Worksheet 3-A(9)(b). Alternatives - Technical F	easibility of Alternatives to Methyl Bromide
In this worksheet, you should address why an alternative pes not effective for your conditions. This worksheet contains 9 of each research study you use to evaluate a single methyl bron	
For worksheet 3-A you must complete one worksheet for each number the worksheets as follows. For the same alternative, same alternative, second research study, label the worksheet the worksheet 3-A(1)(c). For the second alternative, first resealternative, second research study, label the worksheet 3-(A)(	first research study, label the worksheet 3-A(1)(a). For the 3-A(1)(b). For the first alternative, third research study, label arch study, label the worksheet 3-(A)(2)(a). For the second
When completing Section II, if you cite a study that is on the E	EPA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the Resea	rch Summary Worksheet.
If you prefer, you may provide the information requested in th research reports. The narrative review must reply to Section Worksheet of relevant treatments should be provided for each	and questions 1 through 8 in Section II. A Research Summary
BACKGROUND	
EPA must consider whether alternative pest control measures (pessuccessfully instead of methyl bromide by crop and circumstance alternative pest control regimens for various crops, which can be for	
There are three major ways you can provide the Agency with proo	f of your investigative work.
<ul><li>(1) Conduct and submit your own research</li><li>(2) Cite research that has been conducted by others</li><li>(3) Cite research listed on the EPA website</li></ul>	· <u> </u>
Whether you conduct the research yourself or cite studies develop scientifically sound manner. The studies should include a descripti application intervals, pest pressure, weather conditions, varieties of outcome. You must submit copies of each study to EPA unless	on of the experimental methodology used, such as application rates, f the crop used, etc. All results should be included, regardless of
The Agency has posted many research studies on a variety of crop EPA will add studies to its website as they become publicly available websites for studies that pertain to your crop and geographic area.	ole. You are encouraged to review the EPA website and other
In addition, EPA acknowledges that, for certain circumstances, sor research has been conducted (i.e. solarization may not be feasible the Agency and explain why they cannot be used for your crop and	in Seattle). You should look at the list of alternatives provided by
Use additional p	ages as needed.
	ynamics of mefenoxam insensitivity in a recombining ation of <i>Phytophthora capsici</i> characterized with
	ied fragment length polymorphism markers.
Section I. Initial Screening on Technical Fe	easibility of Alternatives
1. Are there any location-specific restrictions that inhibit the u	se of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

1d. Other (Please describe)

For EPA Use Only	
ID#	

## Worksheet 3-A(9)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. K.H. Lamour 2. Author(s) or researcher(s) M.K. Hausbeck Phytopathology 91:553-557, 2001 3. Publication and Date of Publication Michigan 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Crop Rotation 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Oospores of the soil-borne fungus play a key role in overwintering and the frequency of mefenoxam insensitivity may not decrease in an agriculturally significant time period (2 years), rendering crop rotation and fungicide use ineffective. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? This study is directly applicable, since it was conducted in Michigan and documents the situation of commercial

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# The Dynamics of Mefenoxam Insensitivity in a Recombining Population of *Phytophthora capsici* Characterized with Amplified Fragment Length Polymorphism Markers

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#### **ABSTRACT**

Lamour, K. H., and Hausbeck, M. K. 2001. The dynamics of mefenoxam insensitivity in a recombining population of *Phytophthora capsici* characterized with amplified fragment length polymorphism markers. Phytopathology 91:553-557.

Recent findings from Michigan suggest that recombination may play a role in the survival and evolution of sensitivity to the fungicide mefenoxam in populations of *Phytophthora capsici* on cucurbit hosts. In 1998, 63 mefenoxam insensitive isolates were recovered from a squash field in which mefenoxam had been applied. Additional isolates were recovered from untreated squash fields planted at this location in 1999 (200 isolates) and the spring of 2000 (34 isolates). Isolates from 1998 and 1999 were characterized using fluorescent amplified fragment length polymorphism (AFLP) markers and all isolates were screened for com-

patibility type and mefenoxam sensitivity. In 1998 and 1999, 92 and 71% of the isolates, respectively, had unique multilocus AFLP genotypes with no identical isolates recovered between years. Seventy-two identical AFLP markers were clearly resolved in both the 1998 and 1999 sample sets, and fixation indices for the 37 polymorphic AFLP loci indicate little differentiation between years. There was no decrease in the frequency of resistant isolates during the 2 years without mefenoxam selection. We conclude that oospores play a key role in overwintering and that the frequency of mefenoxam insensitivity may not decrease in an agriculturally significant time period (2 years) once mefenoxam selection pressure is removed.

Additional keywords: fungicide resistance, genetic diversity, population genetics.

Crown, root, and fruit rot caused by Phytophthora capsici is increasing in Michigan cucurbit production fields, and uninfested land suitable for rotation is becoming increasingly scarce, especially in areas undergoing rapid urban development. The phenylamide fungicide (PAF) mefenoxam is a systemic fungicide that appears to be acting at the level of DNA translation, and is fungistatic to fully sensitive isolates of P. capsici (2,13). Although mefenoxam has been considered by some growers to be helpful, mefenoxam insensitive isolates were reported on bell peppers in North Carolina and New Jersey by Parra and Ristaino in 1998 (18) and have since been recovered from 10 of 11 farms sampled in Michigan (13), as well as, in Georgia (15) and southern Italy (19). Mefenoxam insensitivity in Michigan P. capsici isolates is inherited as a single gene exhibiting incomplete dominance (13), which is consistent with the reports for a variety of other oomycetous organisms (2). Investigations with P. infestans indicate that insensitivity may be conferred by genes at different chromosomal positions (5), suggesting that the basis of insensitivity in different populations may not be identical. Sexual recombination, in particular, has the potential to impact management strategies that employ PAFs because the fully insensitive (two copies of the insensitivity allele) phenotype may be directly generated. P. capsici is heterothallic and the sexual stage is initiated when isolates of opposite compatibility type, designated A1 and A2, come into close association to form thick-walled oospores (4). The asexual stage includes the production of caducous sporangia born on long pedicels, which may release motile zoospores if free water is present. Asexual spores are thought to be responsible for the polyyelic nature of disease development (20).

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PAF resistance in the genus *Phytophthora* and, in particular, the P. infestans-potato pathosystem, is well documented (2,4,9). Until recently, the population structure of P. infestans appeared to be largely clonal outside of P. infestans putative center of origin (6). The recent detection of both P. infestans compatibility types along with increased genotypic diversity in some potato growing regions indicates that the sexual stage is likely active and may significantly impact control strategies that have proved useful in the past (3,8). When PAF resistance in European P. infestans populations increased significantly in the early 1980s, the efficacy of the PAF metalaxyl was only regained after the product was not made available to growers for a period of time (2). This strategy apparently allowed the resistant populations to decline or become extinct and depends on ephemeral populations or, in the case of resident populations, upon a significant cost for resistance outside of selection pressure. A recent study of sensitive versus PAF resistant P. nicotianae isolates from citrus suggests negligible fitness costs for PAF resistance and reports that 2 years without PAF use did not reduce the proportion of resistant isolates in groves (21). Kadish and Cohen report that PAF-resistant P. infestans isolates in Israel were more aggressive in colonizing tuber tissue than sensitive isolates (12).

Novel techniques have been developed recently that allow characterization of DNA-level polymorphism in organisms for which little is known about the genome. An example is the amplified fragment length polymorphism (AFLP) technique introduced by Vos et al. in 1995 (23). This technique relies on restriction enzyme fragmentation of genomic DNA with the concomitant ligation of synthetic adaptors to the DNA fragment ends. Stringent polymerase chain reaction (PCR) amplification using adaptor-complementary primers with additional selective nucleotides allow for the amplification of fragment subsets. DNA fragment subsets are termed fingerprints and may be resolved with a range of techniques (1). AFLP markers have been used on a variety of organ-

isms (14,22) and the procedure generates a large number of reproducible markers (1,22). The limitation that these markers are generally scored as dominant markers (e.g., either present or absent) for diploid organisms requires the use of relatively large sample sets (11,25).

Our null hypotheses are that sexual recombination has a significant impact on the population structure of *P. capsici* in Michigan and that mefenoxam insensitivity may not decrease in the time frame of a typical 2-year rotation outside of mefenoxam selection pressure.

### MATERIALS AND METHODS

Field plot. Research was conducted on a commercial farm in southwest Michigan, with a history (>11 years) of *P. capsici* on bell peppers and squash and intensive use of PAF. The 4.05-ha field sampled had previously been cropped to soybeans and corn with no known record of *P. capsici* susceptible crops (e.g., tomatoes, peppers, or cucurbits) prior to 1997. During 1997 and 1998, yellow squash and zucchini grown in this field became diseased with *Phytophthora* crown, root, and fruit rot and the grower applied mefenoxam as part of a disease management strategy (Novartis, Greensboro, NC). In 1998, all isolates recovered were either intermediately or fully insensitive to mefenoxam. Both A1

TABLE 1. Fixation indices  $(F_{ST})$  for 37 amplified fragment length polymorphism loci from unique *Phytophthora capsici* isolates collected from a single Michigan cucurbit field during 1998 (N = 57) and 1999 (N = 141)

Fragmenta	1998 f(aa)b	1999 f(aa)	$F_{ST}^{c}$
45	0.02	0.06	0.018
54	0.29	0.29	0.000
64	0.82	0.55	0.048
104	0.11	0.06	0.007
106	0.11	0.04	0.025
110	0.41	0.36	0.002
130	0.41	0.30	0.009
146	0.47	0.24	0.038
149	0.12	0.27	0.029
154	0.39	0.31	0.004
156	0.53	0.83	0.054
172	0.56	0.33	0.034
189	0.16	0.56	0.121
192	0.16	0.37	0.044
193	0.35	0.20	0.022
211	0.47	0.15	0.088
241	0.48	0.32	0.018
256	0.04	0.01	0.022
258	0.43	0.49	0.002
261	0.55	0.54	0.000
270	0.57	0.41	0.015
282	0.35	0.40	0.002
285	0.51	0.73	.0.030
314	0.51	0.34	0.019
320	0.41	0.51	0.006
333	0.16	0.20	0.002
346	0.36	0.33	0.001
361	0.33	0.49	0.017
383	0.21	0.15	0.005
418	0.40	0.34	0.002
431	0.34	0.32	0.001
438	0.67	0.45	0.028
454	0.65	0.49	0.015
492	0.29	0.40	0.009
504	0.51	0.47	0.001
511	0.38	0.28	0.007
548	0.78	0.78	0.000

<sup>&</sup>lt;sup>a</sup> EcoRI-AC/MseI-CA selectively amplified fragment size in base pairs.

and A2 compatibility types were present, and oospores were detected in diseased fruit. In 1999 and 2000, yellow squash was established in a 1,124-m<sup>2</sup> experimental plot in this field, and mefenoxam was not applied. Diseased plants and fruit were sampled on 20 August 1998 (63 isolates from entire field), June through August 1999 (200 isolates from experimental plot), and 13 July 2000 (34 isolates from experimental plot). All isolates were recovered from single diseased plants or fruit.

Isolate collection and maintenance. Isolation from diseased plant material was made onto BARP (25 ppm of benomyl, 100 ppm of ampicillin, 30 ppm of rifampicin, and 100 ppm of pentachloronitrobenzene)-amended UCV8 (840 ml of distilled water, 163 ml of unclarified V8 juice, 3 g of CaCO<sub>3</sub>, and 16 g of Bacto agar) plates. Procedures for obtaining single zoospore isolates were as previously described (13). Single zoospore cultures were maintained on 30 ppm of rifampicin and 100 ppm of ampicillin (RA)-UCV8 plates and transferred bimonthly. Long-term storage consisted of a single 7-mm plug of expanding mycelium from each single zoospore culture being placed in a 1.5-ml microfuge tube with one sterilized hemp seed and 1 ml of sterile distilled water, incubated for 2 to 3 weeks at 23 to 25°C, and stored at 15°C long term.

Phenotypic characterization. Isolates were screened for compatibility type as previously described (13). Mefenoxam sensitivity was characterized according to the in vitro screening technique described by Lamour and Hausbeck (LH technique) for P. capsici isolates in Michigan (13). Isolates were scored as sensitive (S) if growth on UC-V8 agar amended with 100 ppm of mefenoxam was less than 30% compared with a control, as intermediately sensitive (IS) if between 30 and 90%, and fully insensitive (I) if greater than 90% compared with the unamended control. These mefenoxam sensitivity categories are based on a trimodal distribution of 523 field isolates of P. capsici. Clear modal distributions were only attained when screening was conducted with a single high rate of mefenoxam-amended (100 ppm) media (K. Lamour, unpublished data). These putative mefenoxam sensitivity categories were tested by in vitro crosses (I  $\times$  S, IS  $\times$  IS, IS  $\times$  S, and S × S), and chi-square analysis confirmed that the observed progeny numbers were not significantly different than expected for Mendelian inheritance of an incompletely dominant trait (13).

The LH technique differs from a commonly used method described by Goodwin, Sujkowski, and Fry (GSF technique) (9) for *P. infestans* which uses two levels of amended media (5 and 100 ppm) to differentiate the three mefenoxam sensitivity phenotypes and which has been used to characterize *P. capsici* isolates (15,18,19). Unfortunately, analysis of our in vitro crosses and field isolates by the GSF technique did not resolve a clear modal distribution (K. Lamour, *unpublished data*). Assignment of Michigan *P. capsici* isolates to the S category was the same whether using the LH or GSF technique. The only difference was that some *P. capsici* isolates from Michigan rated as fully insensitive by the GSF technique were rated as intermediately sensitive by the LH technique.

DNA extraction and AFLP fingerprinting. A technique for avoiding bacterial contamination prior to growing isolates for DNA extraction was implemented using a modified Van Teigham cell (4). The uppermost portion of a 7-mm plug of mycelium was placed onto the surface of RA-WA plates (30 ppm of rifampicin, 100 ppm of ampicillin, 1,000 ml of distilled water, and 16 g of Bacto agar) and an autoclaved cap from a 1.5-ml microfuge tube was placed over the plug which forced the isolate to grow through the amended media. Isolates were incubated in the dark for 2 t 3 days before two 7-mm plugs were transferred to approximately 15 ml of RA-UCV8 broth in petri dishes (100 × 15 mm) and incubated in the dark for 3 days at 23 to 25°C. Mycelial mats were washed with distilled water and dried briefly under vacuum before

being frozen to -20°C and lyophilized.

<sup>&</sup>lt;sup>b</sup> Observed frequency of the absent state where "a" represents the absence of a fragment.

 $<sup>^{\</sup>rm c}$   $F_{\rm ST}$  calculated from estimated allele frequencies. According to Wright's qualitative guidelines, values from 0 to 0.05 indicate little genetic differentiation and values from 0.05 to 0.15 indicate moderate genetic differentiation

Lyophilized mats were ground with a sterile mortar and pestle. Whole genomic DNA from approximately 50 mg of ground mycelium was extracted with a plant mini kit (Dneasy: Oiagen Inc., Valencia, CA) according to the manufacturers directions. DNA was quantified (Nucleic Acid QuickSticks; Clontech, Palo Alto, CA) according to the manufacturers directions and approximately 100 ng of DNA was subjected to a restriction/ligation reaction, preselective amplification, and selective amplifications using the PCR core mix, adaptor sequences, core primer sequences, and fluorescent-labeled primers available in an AFLP microbial fingerprinting kit (Perkin-Elmer Applied Biosystems, Foster City, CA, henceforth referred to as PE/ABI) and performed exactly as described in protocol part 402977 Rev A (23). All PCR reactions were performed with a minicycler (MJ Research Inc., Waltham, MA) in 0.2-ml tubes according to the cycling parameters outlined in the microbial fingerprinting protocol.

An initial optimization set of reactions was performed with preselective products from *P. capsici* isolate OP97, which was isolated from a cucumber fruit in 1997 (13). Selective amplifications with the selective primers *EcoRI*-AA, -AC, -AG, and -AT were performed in all 16 combinations with the *MseI*-CA, -CC, -CG, and -CT selective primers. *EcoRI* selective primers, available from PE/ABI, were labeled at the 5' end with either carboxy-fluorescein (FAM), carboxytetramethyrhodamine (TAMRA), or carboxy-4',5'-dichloro-2',7'-dimethoxyfluorescein (JOE) fluorescent dyes. The fluorescent dyes are excited by laser radiation and visualized by their characteristic absorption-emission frequencies. Only the fragments containing an *EcoRI* restriction site are resolved.

Products from three reactions labeled with different colored dyes and a carboxy-X-rhodamine (ROX) size standard were loaded into each lane on a denaturing polyacrylamide gel and the fragments resolved in a DNA sequencer (ABI Prism 377). Results were prepared for analysis in the form of electropherograms using GeneScan Analysis software (PE/ABI). AFLP fragments were scored manually as present (1) or absent (0) using Genotyper (PE/ABI). Only DNA bands that consistently exhibited unambiguous presence or absence profiles were scored.

A single isolate, OP97, was subjected to the aforementioned protocol using three primer pair combinations that were chosen as optimal on three separate occasions, approximately 3 months apart, to test for reproducibility of AFLP profiles.

Clone detection and cluster analysis. AFLP fragments were considered polymorphic if the most common allele was present in less than 95% of the isolates from a given sample set and scored for presence (1) or absence (0) (10). AFLP fragments present in more than 95% of the isolates from a given sample set were considered monomorphic. Analysis of the resulting binary data matrix was performed using NTSYS-pc version 2.02k (Exeter Software, Setauket, NY). Unweighted pair group method with arithmetic averages cluster analysis was performed on the matrix of similarity coefficients calculated from all possible pairwise comparisons of individuals within and among the 1998 and 1999 populations and a tree generated. Isolates showing complete homology at all loci were considered to be clones and except for a single representative isolate were excluded from frequency calculations.

Allele frequency and fixation indices. Allele frequencies for AFLP markers were estimated utilizing the expected relationship between gene and genotype frequencies in a randomly mating population (i.e., Hardy-Weinberg proportions). The frequency of the recessive (absent) allele (q) was calculated from the observed number of recessive homozygote individuals (X) in a sample of n individuals by the formula for dominant markers described by Jorde et al. (11):

$$\hat{q} = \sqrt{x + \frac{1 - x}{4n}}$$

where x = X/n is the observed proportion of individuals that do not display the dominant (present) marker phenotype. In order to test whether the composite genetic profiles from 1998 and 1999 were consistent with a single randomly mating population, the fixation index was calculated for each AFLP loci from the variance in allele frequencies according to the following formula:  $F_{ST} = [(p_1 - p_2)^2/4]/(average p \times average q)$ , where p is the allele frequency for the present state with  $p_1$  and  $p_2$  indicating the two sample populations, and q is the allele frequency for the absent state (10). Fixation indices for individual loci were interpreted according to the qualitative guidelines suggested by Wright (24), where the range 0 to 0.05 indicates little genetic differentiation, range 0.05 to 0.15 indicates moderate genetic differentiation, and greater than 0.25 indicates great genetic differentiation (10).

## RESULTS

AFLP band characterization. Evaluation of the 16 EcoRI + 2-MseI + 2 selective primer pair combinations indicated that EcoRI + AC-MseI + CA gave the most clearly resolved fragment profile and was used to amplify genomic DNA from all isolates in both the 1998 and 1999 sample sets. This primer combination resulted in 72 clearly resolved fragments of which 37 (51%) fragments were polymorphic in both 1998 and 1999 (Table 1). All 72 fragments were present in both 1998 and 1999 and no novel fragments were detected between years. The following 35 fragments (size in base pairs) were monomorphic in both the 1998 and 1999 sample sets: 41, 43, 47, 49, 58, 66, 70, 82, 85, 114, 118, 123, 133, 135, 140, 159, 174, 235, 247, 249, 272, 278, 295, 298, 300, 341, 351, 355, 367, 402, 474, 488, 502, 519, and 527. AFLP profiles for isolate OP97, generated from separate DNA extractions on three separate occasions over a 1-year period, resulted in identical banding patterns with the only difference being minor changes in the intensity of the electropherogram signal. Occasionally individual reactions resulted in poorly resolved fingerprint profiles (e.g., low intensity of signal) and were repeated until signals were deemed optimal.

Phenotypic, genotypic, and gene diversity. No isolates sensitive to mefenoxam were recovered in 1998 or 2000, and single A1 sensitive and A2 sensitive isolates were recovered in 1999 (Table 2). In 1998, 18% of the isolates were intermediately sensitive and 82% were insensitive, in 1999, 2% were sensitive, 28% were intermediately sensitive and 70% were insensitive, and in 2000, 15% of the isolates were intermediately sensitive and 85% were insensitive to mefenoxam (Table 2).

In 1998, 57 of the 63 isolates recovered, and 141 of the 200 isolates recovered in 1999 were unique based on multilocus AFLP profiles. No identical multilocus genotypes were recovered between 1998 and 1999. Five isolates (two A2/I, two A2/IS, and

TABLE 2. Phenotypic diversity of *Phytophthora capsici* isolates recovered from the same cucurbit field in 1998, 1999, and 2000

	No. of	Compatibility type and mefenoxam sensitivity <sup>c</sup>					ityc
Yeara	isolates <sup>b</sup>	A1/S	A1/IS	A1/I	A2/S	A2/IS	A2/I
1998	57	_	4	31	_	6	16
1999	141 .	1(2)	17 (20)	57 (53)	1(1)	23 (18)	42 (47)
2000	34	- ` ´	2 ` ′	8 ′	-`´	3 ` ´	21 `

<sup>&</sup>lt;sup>a</sup> Mefenoxam was applied in 1998 but not in 1999 or 2000.

b Sample sets from 1998 and 1999 consist of unique multilocus genotypes as determined with amplified fragment length polymorphism fingerprinting. The 2000 sample set was recovered at the beginning of the growing season and was not fingerprinted.

c S = sensitive, IS = intermediately sensitive, and I = insensitive as determined by in vitro screening on 100 ppm of mefenoxam-amended agar. Numbers in parentheses indicate the expected number of isolates when mefenoxam insensitivity is assumed to be controlled by a single incompletely dominant gene in Hardy-Weinberg equilibrium unlinked to compatibility type.

one A1/I) of *P. capsici* collected in 1998 had one clonal representative. Fourteen isolates collected in 1999 had between two and four clones (Table 3). A single A1 compatibility type insensitive isolate had 40 clones recovered over the course of the 1999 season and comprised 3% of the early, 15% of the mid-, and 43% of the late sampling intervals (Table 3). The 1999 sampling intervals (early, mid, and late) are based on the dates of sampling and are not intended to reflect stages of plant growth or the epidemiology of *P. capsici*. Cluster analysis of AFLP fingerprint variation indicated no significant clustering of isolates between 1998 and 1999.

The majority (98%) of the 37 polymorphic AFLP fragments showed little genetic differentiation ( $F_{\rm ST} < 0.05$ ) between 1998 and 1999 according to Wrights qualitative criterion (Table 1) (24).

### DISCUSSION

P. capsici causes significant damage to cucurbit hosts in Michigan each year. In an effort to prevent or control epidemics, many growers have used either metalaxyl or the newer, but similarly acting compound, mefenoxam as a part of their disease management strategy. This study was initiated in an effort to address the concerns of growers who have high levels of mefenoxam insensitivity.

Phenotypic data (mefenoxam sensitivity and compatibility type) from a 1998 survey suggested that insensitivity to mefenoxam was common and that some level of recombination is occurring in the field (13), but without the application of additional polymorphic markers our ability to assess population structure was severely restricted. AFLP analysis proved to be a powerful tool for resolving the population dynamics of *P. capsici*. A single selective primer combination, *EcoRI-AC-MseI-CA*, generated 72 bands of which 37 were polymorphic in our 1998 and 1999 sample sets. AFLP fingerprinting, in conjunction with temporal sampling, provided a useful characterization of *P. capsici* from one season to the next and allowed us to track asexual disease development over the course of a single season.

Our data suggests that sexual recombination significantly impacts the structure of this *P. capsici* population. The finding that 198 of the 262 isolates recovered between 1998 and 1999 had unique multilocus AFLP genotypes is consistent with the high level of genotypic diversity expected in an outcrossing population

TABLE 3. Clone contribution of 15 *Phytophthora capsici* isolates to the total number of isolates collected in 1999 (N = 200)

			` ,		
		•	No. of clones	in early, mid, ar	nd late season <sup>c</sup>
Isolate	No. of clones	CT/MSb	6/22 - 7/16 N = 60	7/20 - 8/3 N = 80	8/5 - 8/18 N = 60
JP571	2	A1/I	2	_	_
JP583	2	A1/I	2	_	_
JP944	3	A1/I	2	1	
JP999	3	A1/I	2	1	_
JP1007	2	A1/I	1	1	_
JP1042	2	A2/I	1	1	_
JP1096	2	A1/I	_	1	1
JP1102	2	A2/I	-	2	_
JP1215	3	A2/I	3	_	_
JP1342	2	A2/IS	_	2	-
JP1369	2	A 1/I	1	1	_
JP1384	4	A2/I	3	1	-
JP1512	2	A1/I	1	_	1
JP1555	3	A1/I	_		3
JP1632	40	A1/I	2	12	26

<sup>&</sup>lt;sup>a</sup> Total number of isolates with identical multilocus amplified fragment length polymorphism profiles.

(7,16,17). Although clonal reproduction occurred in 1998 and 1999, no identical genotypes were recovered between years, suggesting that oospores are important for overwintering. The finding that 35 of the 37 polymorphic fragments exhibited very little differentiation (i.e., change in allele frequency) based on the estimated fixation indices between 1998 and 1999 is consistent with the expectations for a recombining population large enough to avoid dramatic changes due to genetic drift.

In 1999 and 2000, sensitive and intermediately sensitive isolates (42 of 175) did not increase in a manner suggesting selection in favor of mefenoxam sensitivity outside of mefenoxam selection pressure. The fact that 14 of the 15 isolates with clonal reproduction in 1999 were fully insensitive may be another indication that mefenoxam insensitivity does not have significant costs outside of mefenoxam selection pressure. If we assume that there is only a single mefenoxam insensitivity gene in this population unlinked to compatibility type, designated I, and that this population is effectively free from the effects of migration and genetic drift, some interesting speculations can be made. For instance, in 1999, if the mefenoxam sensitivity phenotypes are assumed to represent genotypes (e.g., a fully insensitive isolate has two copies of the I allele) then the frequency of I can be estimated and the observed number of unique isolates that fall into each of the six mefenoxam sensitivity/compatibility type categories can be compared with the expectations under Hardy-Weinberg equilibrium. In 1999, the estimated frequency of I was 0.84, and chi-square analysis, using the data in Table 2, indicates that the observed numbers do not differ from those expected under Hardy-Weinberg equilibria at  $P = 0.50 \ (\chi^2 = 3.09, \ df = 4)$ . Although this is not a particularly powerful test due to the large number of assumptions (10), it does lend support to the hypothesis that this population meets the criterion for panmixia.

Our results do not allow us to reject the null hypothesis that sexual recombination significantly impacts the structure of this population. It appears that sexual recombination plays a significant role in maintaining genotypic and gene diversity while concomitantly producing overwintering inoculum. Our data also suggest that sexual recombination may serve as a potent force for integrating a beneficial allele based on the finding that there were a total of 133 unique multilocus genotypes fully insensitive to mefenoxam between 1998 and 1999. An interesting question that can only be answered by following a fully sensitive population as it shifts to insensitivity is how much genetic diversity is lost, if any, during the PAF selection process? The question of how long mefenoxam resistance will remain in a population of P. capsici when selection pressure is removed can only be answered in a tentative way. It appears that in this population, insensitivity will not decrease within the time frame of a typical 2-year rotation and, once resistance to mefenoxam is established, the future usefulness of this fungicide may be extremely limited.

Comparison of the population structure reported at this single location is currently being compared with other locations in Michigan and the United States and should provide useful insight into the amount of genetic diversity in sensitive versus insensitive populations as well as the contribution of migration to *P. capsici* population structure.

### **ACKNOWLEDGMENTS**

This work was funded by the Michigan Agricultural Experiment Station, Michigan State University Extension, Michigan Department of Agriculture, Michigan Farm Bureau (GREEEN cooperative), Pickle and Pepper Research Committee, Pickle Packers International Inc., and the Pickle Seed Research Fund, Pickle Packers International. We thank A. M. Jarosz for comments on the manuscript and valuable criticism during this project, E. A. Webster for supervision of lab procedures, and M. Bour, C. Hunter, J. Jabara, and P. Tumbalam for competent lab assistance.

CT = compatibility type and MS = mefenoxam sensitivity where S = sensitive, IS = intermediately sensitive, and I = insensitive as determined by in vitro screening on 100 ppm of mefenoxam-amended agar.

<sup>&</sup>lt;sup>c</sup> Sample intervals based on sampling dates only.

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## Worksheet 3-A(9)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Crop Rotation, Fallow
Study: Investigating the spatiotemporal genetic structure of
Phytophthora capsici in Michigan.

## Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the	use of this alternative	e on your site?	
1a. Full use permitted	X		
1b. Township caps			
1c. Alternative not acceptable in consuming country			
1d. Other (Please describe)			

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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## Worksheet 3-A(9)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) K.H. Lamour M.K. Hausbeck 3. Publication and Date of Publication Phytopathology 91:973-980, 2001 4. Location of research study Michigan, USA 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Crop Rotation 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Crop rotation is not highly effective because both mating types of *Phytophthora capsic* i are present in Michigan fields, resulting in an oospore capable of surviving for long period of time. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Results are directly applicable, since the research was conducted in Michigan.

OMB Control # 2060-0482

# Investigating the Spatiotemporal Genetic Structure of *Phytophthora capsici* in Michigan

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#### **ABSTRACT**

Lamour, K. H., and Hausbeck, M. K. 2001. Investigating the spatiotemporal genetic structure of *Phytophthora capsici* in Michigan. Phytopathology 91:973-980.

Phytophthora capsici isolates were recovered from pepper and cucurbit hosts at seven locations in Michigan from 1998 to 2000. Isolates were characterized for compatibility type (CT), mefenoxam sensitivity (MS), and amplified fragment length polymorphism (AFLP) marker profiles. In total, 94 AFLP bands were resolved. Individual populations were highly variable. Within populations, 39 to 49% of the AFLP bands were polymorphic and estimated heterozygosities ranged from 0.16 to 0.19. Of the 646 isolates fingerprinted, 70% (454) had unique AFLP

profiles. No clones were recovered between years or locations. Pairwise F statistics ( $\Phi_{ST}$ ) between populations from different locations ranged from 0.18 to 0.40. A tree based on unweighted pair-group method with arithmetic average cluster analysis indicates discrete clusters based on location. Isolates from the same location showed no clustering based on the year of sampling. Analysis of molecular variance partitioned variability among (40%) and within populations (60%). The overall estimated  $\Phi_{ST}$  was 0.34 (SD = 0.03). A1/A2 CT ratios were  $\approx$ 1:1, and MS frequencies were similar between years for the two locations sampled over time. These data suggest that P. capsici persists in discrete outcrossing populations and that gene flow among locations in Michigan is infrequent.

Phytophthora capsici Leonian causes significant damage to a variety of plant hosts worldwide, and in the United States, it seriously impacts the production of cucurbits, tomatoes, and peppers (9,14,20). In Michigan, the life history of P. capsici is divided between an active growth phase in the presence of susceptible host tissue and a state of dormancy over the winter. Overwintering survival is thought to be accomplished by thick-walled oospores that are produced during sexual reproduction (9.10). P. capsici is heterothallic, and completion of the sexual stage requires both A1 and A2 compatibility types (CT). Sexual reproduction is mediated by extracellular hormonal signals, and there is the potential for both self and cross-fertilization (8). Oospores generally require a dormancy period prior to germination. Germinating oospores produce coenocytic mycelium, which can directly infect or differentiate into caducous sporangia under suitable conditions. Sporangia can be dislodged and cause infection directly, or, in the presence of free water, release 20 to 40 motile zoospores. Polycyclic asexual spread of P. capsici between and down rows has been clearly documented in the pepper/P. capsici pathosystem (21).

Ristaino and Johnston recently summarized management strategies useful for disease control (20). The primary strategy is to manage soil water dynamics by providing the best possible drainage for the host plant's rhizosphere and the field, in general. Growers are advised to rotate fields to nonsusceptible hosts, and when appropriate to apply fungicides.

The phenylamide fungicide mefenoxam is fungistatic to sensitive isolates of *P. capsici* (16), but, as has occurred with many oomycetes, insensitivity has developed in field populations (9,17,18). Research with *P. capsici* isolates from Michigan indicates that insensitivity is controlled by an incompletely dominant

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gene of major effect (9), which is consistent with the findings for other oomycetes (3).

In Michigan, fruit, stem, and root rots caused by P. capsici on susceptible hosts have increased in recent years, and growers employing available management strategies have experienced significant losses. Over the last 4 years, an investigation of P. capsici populations in Michigan commercial vegetable production fields has been conducted (9,10). The initial phase of this study was based on the distribution and frequency of CT and mefenoxam sensitivity (MS) phenotypes within fields. In 1998, an approximate 1:1 ratio of A1/A2 isolates was discovered in the majority of fields sampled, and oospores were detected in diseased cucurbit fruit on four separate farms. All six CT/MS phenotypes were recovered as oospore progeny from a single diseased cucumber fruit as well as from a single diseased cucumber field (9). These initial findings suggested that the sexual stage occurs in populations of P. capsici in Michigan and, based on the MS findings, that sexual recombination may play an important role in generating the fully insensitive MS phenotype.

The ability to assess population structure by only CT and MS is limited by the fact that only six phenotypic combinations are resolvable and is further limited because some populations appear to have only sensitive or insensitive isolates (9). Amplified fragment length polymorphism (AFLP) markers are increasingly used as a tool to investigate population genetic structure in a wide variety of living organisms including plants (22,24), animals (19), insects (4), and microorganisms (11). A molecular map of the P. infestans genome was constructed based on AFLP and restriction fragment length polymorphism markers and corroborates the finding of other researchers that AFLP markers span the genome (23). The recent characterization of a single mefenoxam-insensitive population of P. capsici with AFLP markers over a 2-year period revealed that genotypic and genic diversity were high, that clonal reproduction (the recovery of identical multilocus genotypes from different locations within a field) was significant within a single season but that members of the same clonal lineage were not detected between years, that AFLP marker frequencies did not

change significantly between years, and that the frequency of mefenoxam insensitivity did not appear to decrease in the absence of mefenoxam use (10).

In this paper, we report on the genetic structure of *P. capsici* populations from fields located in different regions of Michigan. It was our goal to consider dispersal between locations and the impact of outcrossing on natural populations. We also report on the frequency of self-fertilized versus outcrossed progeny in a sexual cross between isolates from different geographical locations and the inheritance of AFLP markers in this cross. Portions of the information in this paper have been reported previously (9,10).

### MATERIALS AND METHODS

Isolate collection and maintenance. Pepper, cucumber, pumpkin, tomato, and squash plant tissue (root, crown, and fruit) with typical signs and symptoms of infection by P. capsici were collected from six farms in four different regions of Michigan between 1998 and 2000. Sampling was conducted using fieldspecific grids with grid quadrants varying from 40 m<sup>2</sup> to 12 km<sup>2</sup>, depending on the size of the field. Sample sets are labeled according to the following notational approach: location (SW = southwest, SC = south central, C = central, and NW = northwest) followed by a farm designation (1,2,...n) with a hyphen separating a field designation (A,B,...n) and the year sampling was conducted (98, 99, and 00). Diseased plant tissue (between 4 to 20 per quadrant) was collected from quadrants in an arbitrary fashion. Isolation from diseased plant material was made onto BARP (25 ppm of benomyl, 100 ppm of ampicillin, 30 ppm of rifampicin, and 100 ppm of pentachloronitrobenzene)-amended UCV8 (840 ml of distilled water, 163 ml of unclarified V8 juice, 3 g of CaCO<sub>3</sub>, and 16 g of agar) plates. Procedures for obtaining single zoospore isolates were as previously described (9). Single zoospore cultures were maintained on RA (30 ppm of rifampicin and 100 ppm of ampicillin)-UCV8 plates and transferred bimonthly. For long-term storage, a 7-mm plug of expanding mycelium from each culture was placed in a 1.5-ml microfuge tube with one sterilized hemp

TABLE 1. Inheritance of 17 amplified fragment length polymorphism (AFLP) markers, compatibility type (CT), and mefenoxam sensitivity (MS) in 107 progeny of a cross between *Phytophthora capsici* isolates OP97 (A1/IS) and SFF3 (A2/S)

(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	<u></u>		
Marker <sup>a</sup>	Progeny ratiob	χ <sup>2c</sup>	P <sup>d</sup>
E+AC/M+CA-66	47:60	1.58	0.20
E+AC/M+CA-97	51:56	0.23	0.70
E+AC/M+CA-146	53:54	0.01	0.90
E+AC/M+CA-149	60:47	1.58	0.20
E+AC/M+CA-156	64:43	4.12	0.04
E+AC/M+CA-159	56:51	0.23	0.70
E+AC/M+CA-244	46:61	2.10	0.17
E+AC/M+CA-258	52:55	0.08	0.80
E+AC/M+CA-270	53:54	0.01	0.98
E+AC/M+CA-282	56:51	0.23	0.70
E+AC/M+CA-290	62:45	2.70	0.13
E+AC/M+CA-328	55:52	0.08	0.80
E+AC/M+CA-351	61:46	2.10	0.15
E+AC/M+CA-398	55:52	0.08	0.80
E+AC/M+CA-431	55:52	0.08	0.80
E+AC/M+CA-435	57:50	0.46	0.90
E+AC/M+CA-444	49:58	0.76	0.85
CT	53:54	0.01	0.98
MS	47:60	1.58	0.20

AFLP marker labels indicate the restriction enzymes (E = EcoRI, M = MseI), the two selective nucleotides, and the size of the DNA fragment in base pairs.

seed and 1 ml of sterile distilled water (SDW). Isolates were then incubated for 2 to 3 weeks at 23 to 25°C before being stored at 15°C.

CT and MS determination. Agar plugs from the edge of an expanding single zoospore colony were placed at the center of UCV8 plates approximately 2 cm from ATCC isolate 15427 (A1 CT) and ATCC 15399 (A2 CT) and incubated in the dark at 23 to 25°C for 3 to 6 days. Following incubation, CT was determined. Thereafter, all CT determinations were crossed with field isolates OP97 (A1) and SP98 (A2).

Agar plugs from the edge of actively expanding single zoospore colonies were placed at the center of UCV8 plates (100 × 15 mm) amended with 0 or 100 ppm of mefenoxam (Ridomil Gold EC, Novartis, Greensboro, NC; 48% active ingredient, suspended in SDW; added to UCV8 cooled to 49°C). Inoculated plates were incubated at 23 to 25°C for 3 days, and colony diameters were measured. Percent growth of an isolate on amended media was calculated by subtracting the inoculation plug diameter (7 mm) from the diameter of each colony and dividing the average diameter of the colony on unamended plates by the average diameter of the colony on unamended control plates. All tests were conducted at least two times. An isolate was scored as sensitive (S) if growth at 100 ppm was <30% of the control, intermediately sensitive (IS) if growth was between 30 and 90% of the control, and insensitive (I) if growth was >90% of the control (9).

DNA extraction and AFLP fingerprinting. Bacterial contamination was avoided by using a modified Van Teigham cell (5). The uppermost portion of a 7-mm plug of mycelium was placed on the surface of RA-WA plates (30 ppm of rifampicin, 100 ppm of ampicillin, 1,000 ml of distilled water; and 16 g of agar) and an autoclaved cap from a 1.5-ml microfuge tube was placed over the plug, which forced the isolate to grow through the amended medium. Isolates were incubated in the dark for 2 to 3 days before two 7-mm plugs were transferred to approximately 15 ml of RA-UCV8 broth in petri dishes (100 × 5 mm) and incubated in the dark for 3 days at 23 to 25°C. Mycelial mats were washed with distilled water and dried briefly under vacuum before being frozen to -20°C and lyophilized.

Lyophilized mats were ground with a sterile mortar and pestle. Whole genomic DNA from approximately 50 mg of ground mycelium was extracted with a plant mini kit (Qiagen Dneasy; Qiagen Inc., Valencia, CA) according to the manufacturer's directions or using a cetyltrimethylammonium bromide (CTAB) procedure in conjunction with an automated DNA extractor. DNA was

TABLE 2. Estimates of genetic diversity within populations of *Phytophthora capsici* in Michigan

Population <sup>a</sup>	No. of isolates <sup>b</sup>	No. of AFLP bands	No. and % polymorphic bands <sup>c</sup>	Estimated average heterozygosity
SW1-A98	57	72	37 (39)	0.16
SW1-A99	141	72	37 (39)	0.16
SW1-B99	35	69	38 (40)	0.16
SW1-B00	24	69	38 (40)	0.16
SC1-A98	50	68	42 (45)	0.17
SC2-B99	45	71	43 (46)	0.17
C1-A00	48	77	41 (44)	0.17
NW1-A99	37	80	44 (47)	0.19
NW2-B98	24	73	46 (49)	0.18

<sup>&</sup>lt;sup>a</sup> First two capital letters indicate location in Michigan with S = south, W = west, C = central, and N = north, the number following the location designator indicates the farm, the capital letter following the hyphen is a field designator, and the numbers following the field designator indicate year (e.g., 00 = 2000).

b Presence/absence ratios for AFLP markers, A1/A2 for CT, and sensitive (S)/ intermediately sensitive (IS) for MS as determined by in vitro screening.

 $<sup>^{</sup>c}$   $\chi^{2}$  value for testing 1:1 segregation (1 df).

<sup>&</sup>lt;sup>d</sup> Probability of the observed ratio occurring by chance under the null hypothesis of 1:1 segregation.

b Total number of isolates with unique multilocus amplified fragment length polymorphism (AFLP) profiles.

e Percent polymorphic bands determined by dividing the number of polymorphic bands by the total number of bands recovered in Michigan (N = 94).

quantified by Nucleic Acid QuickSticks (Clontech, Palo Alto, CA) according to the manufacturer's directions or on 1.5% agarose gels. Approximately 100 ng of DNA was subjected to a restriction/ligation reaction, preselective amplification, and selective amplifications using the polymerase chain reaction (PCR) core mix, adaptor sequences, core primer sequences, and fluorescence-labeled primers provided by an AFLP microbial fingerprinting kit (Perkin-Elmer Applied Biosystems [PE/ABI], Foster City, CA) and performed exactly as described in the AFLP microbial finger-printing protocol part 402977 Rev A (PE/ABI) (25). All PCR reactions were performed with a minicycler (MJ Research Inc., Waltham, MA) in 0.2-ml tubes according to the cycling parameters outlined in the microbial fingerprinting protocol.

An initial optimization set of reactions was performed with preselective products from *P. capsici* isolate OP97, which was isolated from a cucumber fruit in 1997 (9). Selective amplifications with the selective primers *EcoRI*-AA, AC, AG, and AT were performed in all 16 combinations with the *MseI*-CA, CC, CG, and CT selective primers. *EcoRI*-selective primers available from PE/ABI were labeled at the 5' end with either carboxyfluorescein (FAM), carboxytetramethyrhodamine (TAMRA), or carboxy-4',5'-dichloro-2',7'-dimethoxyfluorescein (JOE) fluorescent dyes. The fluorescent dyes were excited by laser radiation and visualized by their characteristic absorption-emission frequencies. Only the fragments containing an *EcoRI* restriction site were resolved.

Selective amplification AFLP products and a carboxy-X-rhodamine size standard were loaded into each lane on a denaturing polyacrylamide gel and the fragments resolved in a DNA sequencer (Prism 377; ABI). Results were prepared for analysis in the form of electropherograms using GeneScan Analysis software (PE/ABI). AFLP fragments were scored manually as present = 1 or absent = 0 using Genotyper (PE/ABI). Only DNA bands that consistently exhibited unambiguous presence/absence profiles were scored.

In order to assess the reproducibility of AFLP profiles, a single isolate, OP97, was subjected to the aforementioned protocol using three optimal primer pair combinations on three separate occasions approximately 3 months apart.

No prior sequencing or cloning of fragments is needed to utilize this marker system and it is highly reproducible between labs (1). AFLP markers are generally scored as present or absent (e.g., dominant markers), and the confidence with which population level inferences can be made is greatly increased by sample sets that are approximately twice the size used for codominant markers (7,12,28).

Marker inheritance. Oospore progeny (N=107) resulting from a cross between isolate OP97 (A1/IS) × SFF3 (A2/S) were subjected to AFLP analysis as described previously. Protocols for the generation, germination, and phenotypic characterization of the F1 oospores from this cross have been reported previously (9). The inheritance of AFLP bands present in one parent and absent in the other were analyzed by chi-square analysis to compare observed numbers to those expected under simple Mendelian inheritance (23). Bands present in a single parent and inherited in all the progeny were assumed to be present in two copies in the parent. Bands present in both parents, or present in two copies in one parent and absent in the other, are not reported on in this study. Individual oospore isolates were evaluated to determine if they were the products of self-fertilization or outcrossing between the parent isolates.

Clone detection. AFLP fragments for each field isolate were scored for presence or absence, and the binary data matrix was converted to a similarity matrix using a simple matching coefficient of resemblance with the program NTSYS-pc version 2.02k (Exeter Software, Setauket, NY). Unweighted pair group method with arithmetic averages (UPGMA) cluster analysis was performed on the similarity matrix and a tree was generated. Isolates showing complete homology at all loci were considered members of the same clonal lineage and, except for a single representative

isolate (referred to as a clone), were excluded from population genetic analysis (13).

Population genetic analysis. Sample sets collected from single fields during a single year were considered a population. Populations were assumed to be in Hardy-Weinberg equilibrium, and each AFLP locus was assumed to be diallelic and selectively neutral. The program tools for population genetic analysis (TFPGA) (M. P. Miller, Northern Arizona University, Flagstaff) was used to assess genetic diversity within each population on the basis of estimated average heterozygosity (15) and the proportion of polymorphic loci at the 95% level (6), and to calculate pairwise and overall fixation indices (F statistics) according to the methods of Weir and Cockerham (26). Confidence intervals for F statistics at the 95% confidence level were generated by bootstrapping using 1,000 iterations.

The fixation index, as described by Wright, equals the reduction in heterozygosity expected with random mating at any one level of a population hierarchy relative to another more inclusive level of the hierarchy (27). Weir and Cockerham's approach to estimating fixation indices attempts to correct for the effects of sampling a limited number of organisms from a limited number of populations and is reported as  $\Phi_{ST}$  instead of  $F_{ST}$  (26). Theoretically, the fixation index has a minimum of 0 (no loss of heterozygosity between the populations compared) and a maximum of 1 (indicating fixation for alternative alleles in different populations or a total loss of heterozygosity), but, as discussed by Hartl and Clark (6), the observed maximum is usually much less than 1. Wright (27) has suggested the following qualitative guidelines for the interpretation of fixation indices: the range 0 to 0.05 indicates little genetic differentiation, 0.05 to 0.15 indicates moderate genetic differentiation, 0.15 to 0.25 indicates great genetic differentiation, and values above 0.25 indicate very great genetic differentiation.

Using the program NTSYS-pc, the combined 0/1 data matrix for isolates from all populations was used to construct a genetic similarity matrix of all possible pairwise comparisons of individuals within and among populations using Jaccard's similarity coefficient: GS(ij) = a/(a + b + c). GS(ij) is the measure of genetic similarity between individuals i and j, where a is the number of polymorphic bands shared by i and j, b is the number of bands present in i and absent in j, and c is the number of bands present in j but absent in i. Trees were constructed using UPGMA cluster analysis to provide a graphic representation of the relationships among isolates. A cophenetic correlation coefficient was computed to assess the goodness of fit of the tree to the similarity matrix.

TABLE 3. Clonal component of genotypic diversity within populations of *Phytophthora capsici* from Michigan

	£ 6						
Population <sup>a</sup>	Total no. of isolates	Unique AFLP genotypes (%) <sup>b</sup>	No. of clonal lineages	Minimum:maximum no. of isolates per clonal lineage			
SW1-A98	63	57 (90)	5	2:2			
SW1-A99	200	141 (71)	15	2:40			
SW1-B99	71	34 (48)	12	2:9			
SW1-B00	36	24 (67)	5	2:8			
SC1-A98	57	50 (88)	5	2:3			
SC2-B99	56	45 (80)	5	2:5			
C1-A00	51	48 (94)	3	2:2			
NW1-A99	88	37 (42)	12	2:12			
NW2-B98	24	18 (75)	3	2:3			
Total	646	454 (70)	65	•••			

<sup>&</sup>lt;sup>a</sup> First two capital letters indicate location in Michigan with S = south, W = west, C = central, and N = north, the number following the location designator indicates the farm, the capital letter following the hyphen is a field designator, and the numbers following the field designator indicate year (e.g., 0 = 2000).

b Percentages calculated by dividing the number of unique amplified fragment length polymorphism (AFLP) genotypes by the total number of isolates recovered Genetic structure was also examined by analysis of molecular variance (AMOVA) using the ARLEQUIN software package (L. Excoffier, University of Geneva). The AMOVA analysis was used to partition the variance in banding patterns within and among populations from the same geographical site over consecutive years, between sites on the same farm separated by approximately 1 km, and between all the locations sampled in Michigan. Significance values were assigned to variance components based on a set of null distributions generated by a permutation process, which randomly assigned individuals to populations and drew 1,000 independent samples. In order to clearly summarize the spatial aspect of genetic differentiation, regression was used to fit an appropriate model to the plot of pairwise  $\Phi_{ST}$  values and geographical distances.

### RESULTS

AFLP band characterization and marker inheritance. Evaluation of 16 EcoRI + 2/MseI + 2 selective primer pair combinations indicated that EcoRI + AC/MseI + CA (EAC/MCA) provided the most clearly resolved fragment profile, and this primer pair was used to analyze DNA from the isolates in this investigation. AFLP profiles for isolate OP97, generated from separate DNA extractions on three separate occasions over a 1-year period, were identical, with only minor differences in the intensity of the electropherogram signal. Occasionally, individual reactions resulted in poorly resolved fingerprint profiles (e.g., low intensity of signal) and were repeated until signals were deemed optimal. The EAC/MCA primer combination resulted in 94 clearly resolved fragments between 40 and 550 bps when considering all the isolates recovered from Michigan.

AFLP analysis of oospore progeny from cross OP97 × SFF3 revealed that all 107 progeny had a combination of bands that were present in only a single parent, indicating that each was a product of outcrossing and not self-fertilization. A comparison of the observed to the expected ratios (1:1) for 17 bands, which were present in only one parent, indicated that only one band segregated in a manner significantly different than expected (P = 0.05) (Table 1). Chi-square analysis also indicated that the observed ratios of A1/A2 CT and S/IS MS were not significantly different than expected under Mendelian inheritance (Table 1).

Gene and genotypic diversity. Each isolate was scored for the presence or absence of all 94 AFLP bands. The number of AFLP bands present in each population ranged from 68 to 80, with an average of 72; the number of polymorphic bands ranged from 39 to 49, with an average of 43; and the estimated average heterozygosity ranged from 0.16 to 0.19, with an average of 0.17 (Table 2). These measurements fall within the range described for a wide range of obligately outcrossing diploid plant species. Seventeen (18%) AFLP loci were fixed for the present state (every isolate analyzed had these AFLP markers) in all populations; 12 (13%) were polymorphic in all populations, and 65 (69%) were fixed for presence or absence in some populations and polymorphic in others. The high proportion of AFLP markers fixed among the populations gives a strong indication that significant genetic differentiation exists.

Of the 646 isolates analyzed, 70% had unique multilocus AFLP profiles (Table 3). This suggests that inoculum originating from oospores plays a surprisingly large role in contributing to epidemic development. The number of clonal lineages detected from single locations in Michigan varied from 3 to 15, and the number of isolates within any single clonal lineage ranged from 2 to 40

TABLE 4. Pairwise F statistics  $(\Phi_{ST})^a$  (below diagonal) and geographical distances (in kilometers, above diagonal) among *Phytophthora capsici* populations in Michigan

Populations <sup>b</sup>	SW1-A98	SW1-A99	SW1-B99	SW1-B00	SC1-A98	SC2-B99	C1-A00	NW1-A99	NW2-B98
SW1-A98		0	1	1	165	169	150	180	185
SW1-A99	0.04		1	1	165	169	150	180	185
SW1-B99	0.18	0.25		0	166	170	150	180	185
SW1-B00	0.25	0.24	0.03	•••	166	170	150	180	185
SC1-A98	0.36	0.37	0.29	0.29	•••	8	135	260	265
SC2-B99	0.33	0.35	0.32	0.33	0.28	•••	130	255	260
C1-A00	0.36	0.37	0.33	0.32	0.38	0.40		140	145
NW1-A99	0.32	0.34	0.30	0.30	0.32	0.32	0.38		5
NW2-B98	0.36	0.37	0.31	0.32	0.33	0.33	0.33	0.27	

<sup>&</sup>lt;sup>a</sup> Estimated fixation index calculated according to the methods of Weir and Cockerham.

TABLE 5. Results of nested analysis of molecular variance (AMOVA) for Phytophthora capsici isolates based on 94 amplified fragment length polymorphism markers

Source of variation <sup>a</sup>	Degrees of freedom	Sum of squares	Variance component	Percent variation	P <sup>b</sup>
SW1-A (1998-1999)					
Among populations	1	39.658	0.396	5.05	< 0.0001
Within populations	197	1,461.559	7.457	94.95	
SW1-B (1999-2000)					
Among populations	1	6.678	0.016	0.27	0.0029
Within populations	57	312.399	6.248	99.73	
SW1-A vs. SW1-B					
Among populations	1	234.790	2.762	27.34	< 0.0001
Within populations	255	1,820.294	7.340	72.66	
All locations					
Among populations	6	1,169.295	4.814	39.67	< 0.0001
Within populations	273	1,984.345	7.322	60.33	

<sup>&</sup>lt;sup>a</sup> First two capital letters indicate location in Michigan with S = south, W = west, C = central, and N = north, the number following the location designator indicates the farm, and the capital letter following the hyphen is a field designator. Variance is partitioned between 1998 and 1999 at SW1-A, between 1999 and 2000 at SW1-B, between combined sample sets from SW1-A and SW1-B, and within and between sample sets from seven locations in Michigan. AMOVA analysis for all locations includes sample sets from a single year for locations SW1-A and SW1-B.

b First two capital letters indicate location in Michigan with S = south, W = west, C = central, and N = north, the number following the location designator indicates the farm, the capital letter following the hyphen is a field designator, and the numbers following the field designator indicate year (e.g., 00 = 2000).

b P = the probability of obtaining a more extreme variance component estimate by chance alone based on 1,000 sampling realizations.

(Table 3). In all cases, isolates with identical multilocus AFLP profiles had identical CT and fell into the same MS category. No clones were recovered among separate locations, and cluster analysis indicated that isolates from the same location grouped discretely. The percentage of genotypically unique isolates recovered at locations ranged between 42 and 94% (Table 3). This wide variation may be due to when the samples were collected. Sample sets collected at SW1-A over the course of the 1999 growing season exhibited significantly less genotypic diversity at the end of the season due to the spread of clonal lineages (10). This is expected for an organism with polycyclic disease develop-

ment and suggests that samples collected early in a *P. capsici* epidemic may provide a better estimate of genic diversity than samples collected at the height of an epidemic.

Temporal dynamics. F statistics ( $\Phi_{ST}$ ) comparing populations of P. capsici recovered from field SW1-A over 1998 and 1999, and field SW1-B over 1999 and 2000 were 0.04 and 0.03, respectively. These values indicate that very little genetic differentiation or loss of heterozygosity occurred between years at either location (Table 4). At both locations, the number and identity of AFLP bands resolved remained identical over time, with 72 total bands recovered from populations at SW1-A and 69 bands recovered

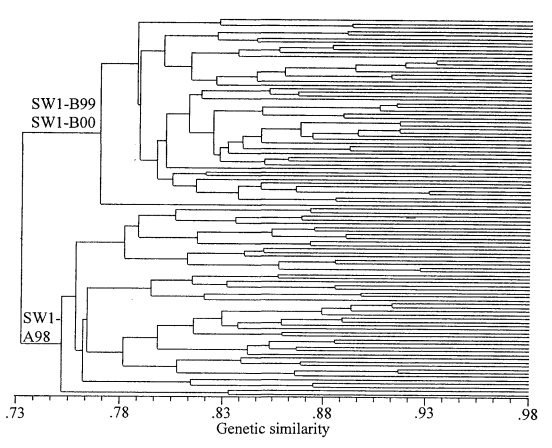


Fig. 1. Unweighted pair-group method with arithmetic average cluster analysis of *Phytophthora capsici* isolates from location SW1-B over 1999 and 2000 (N = 58) and SW1-A in 1998 (N = 57) based on the Jaccard similarity coefficient using 94 amplified fragment length polymorphism markers. Nodes contain isolates exclusively from single locations. Location identifiers precede the inclusive node and are indicated by region (S = 500) south, S = 5000 morth, S = 5000 mort

TABLE 6. Location, year, hosts, compatibility type, and mefenoxam sensitivity of genetically unique Phytophthora capsici isolates collected in Michigan between 1998 and 2000

Population <sup>a</sup>		No. of isolates <sup>c</sup>	Compatibility type/mefenoxam sensitivity <sup>d</sup>					
	Hosts <sup>b</sup>		A1/S	A1/IS	A1/I	A2/S	A2/IS	A2/I
SW1-A98	S, PK	57	•••	4 (0.07)	31 (0.54)		6 (0.11)	16 (0.28)
SW1-A99	S	141	1 (0.01)	17 (0.12)	57 (0.40)	1 (0.01)	23 (0.16)	42 (0.30)
SW1-B99	Ś	34	14 (0.41)	4 (0.12)		11 (0.32)	4 (0.12)	1 (0.03)
SW1-B00	S	24	7 (0.29)	5 (0.21)		5 (0.21)	5 (0.21)	2 (0.08)
SC1-A98	С	50	10 (0.20)	17 (0.34)	2 (0.04)	10 (0.20)	11 (0.22)	2 (0.00)
SC2-B99	С	45		6 (0.13)	22 (0.49)		2 (0.04)	15 (0.33)
C1-A00	P	48	20 (0.42)			28 (0.58)	- (0.0.1)	` '
VW1-A99	S, C	37	25 (0.68)		•••	12 (0.32)		•••
VW2-B98	P	18	10 (0.56)	•••	•••	7 (0.39)	1 (0.05)	•••
<b>Fotal</b>		454	87 (0.19)	53 (0.12)	112 (0.25)	74 (0.16)	52 (0.11)	76 (0.17)

First two capital letters indicate location in Michigan with S = south, W = west, C = central, and N = north, the number following the location designator indicates the farm, the capital letter following the hyphen is a field designator, and the numbers following the field designator indicate year (e.g., 00 = 2000).

<sup>&</sup>lt;sup>b</sup> S = squash, C = cucumber, PK = pumpkin, and P = pepper.

<sup>&</sup>lt;sup>c</sup> Total number of isolates with unique multilocus amplified fragment length polymorphism profiles.

d Mefenoxam sensitivity determined by in vitro screening on 100 ppm of mefenoxam-amended media with S = <30% growth of control (GC), IS = between 30 and 90% GC and I = >90% GC. Observed numbers are followed by proportion of total sample size in parenthesis.

from populations at SW1-B (Table 2). The number and identity of bands polymorphic at the 95% level (37 for SW1-A and 38 for SW1-B) and the estimated average heterozygosity (0.16 for both locations) also remained constant over time (Table 2). AMOVA analysis of SW1-A and SW1-B over time partitioned 5% of the total variability between years for SW1-A and <1% of the total variability between years at SW1-B (Table 5). Significant clonal reproduction was detected at both field sites within a given year, but no members of the same clonal lineage were detected among locations or years (Table 3). Thus, even though individual genotypes did not appear to survive the winter, the data suggest that there was enough outcrossing and survival of the resulting recombinant progeny at both these locations to maintain genic diversity.

Cluster analysis showed that isolates from SW1-A and SW1-B branched from location-specific nodes (branch points on the tree). If there was migration between the locations, then isolates from

SW1-A and SW1-B would be expected to be intermixed in the cluster analysis. On the other hand, there was no clustering within either of the location-specific clusters based on year (Fig. 1). The ratio of A1/A2 CT approximated a 1:1 ratio at each location (Table 6). The percentage of isolates falling into the six MS/CT categories remained relatively similar between years at each location, with a breakdown of 0 and 1% A1/S, 7 and 12% A1/IS, 54 and 40% A1/I, 0 and 1% A2/S, 11 and 16% A2/IS, and 28 and 30% A2/I for location SW1-A in 1998 and 1999, respectively (Table 6). The percentage of isolates in each of the six categories for SW1-B was 41 and 29% A1/S, 12 and 21% A1/IS, 0 and 0% A1/I, 32 and 21% A2/S, 12 and 21% A2/IS, and 3 and 8% A2/I between 1999 and 2000, respectively (Table 6).

Genetic structure. Pairwise  $\Phi_{ST}$  values ranged from 0.18 to 0.40 when comparing populations from different locations (Table 4). According to Wright's criterion, this means that populations

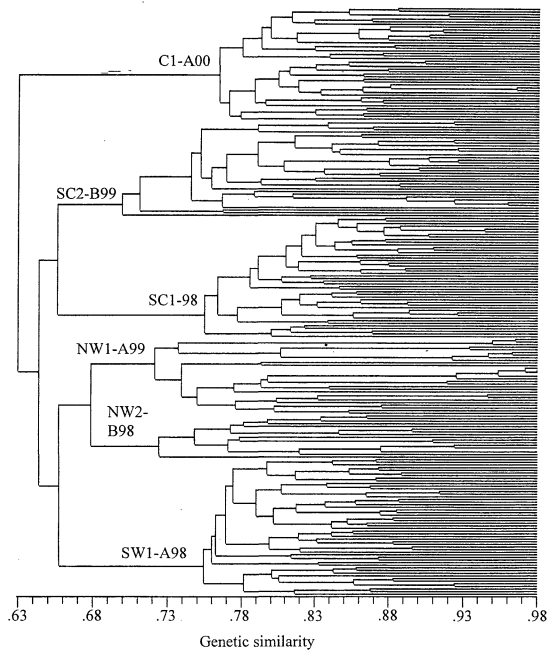


Fig. 2. Unweighted pair-group method with arithmetic average cluster analysis of 255 *Phytophthora capsici* isolates from six locations in Michigan based on the Jaccard similarity coefficient using 94 amplified fragment length polymorphism markers. Nodes contain isolates exclusively from single locations. Location identifiers precede the inclusive node and are indicated by region (S = south, N = north, W = west, and C = central) and a farm identifier (1,2,...n) prior to the hyphen with a field indicator (A, B, ...n) and the year of sampling (e.g., 0 = 2000) following.

were greatly differentiated, even when located as close as 1 km apart. The overall  $\Phi_{ST}$  value when analyzing isolates from all seven locations combined was 0.34 (SD = 0.03), which indicates that approximately 34% of the total genetic variation was present among locations. An AMOVA analysis of sample sets from all locations corroborated this finding and attributed 40% of the genetic variation among populations and 60% within populations (Table 5). Cluster analysis was also in agreement with the overall fixation index and revealed that populations from different geographical locations branched from specific nodes (Figs. 1 and 2), with population-specific clusters being between 63 and 75% similar. Genetic similarities between individuals within each of the clusters showed similar patterns with individuals ranging between 75 to 95% for SW1-A (1998 and 1999), 77 to 94% for SW1-B (1999 and 2000), 75 to 94% for SC1-A98, 69 to 92% for SC2-B99, 76 to 95% for C1-A00, 71 to 97% for NW1-A99, and 72 to 93% similar for NW2-B98 (Figs. 1 and 2). The cophenetic correlation coefficient for the overall tree (Fig. 2) was 0.84, indicating that the tree provided a good fit to the data matrix. The results of fitting a linear model to describe the relationship between pairwise  $\Phi_{ST}$  and pairwise geographical distances indicated a significant relationship ( $r^2 = 42.67$ ; P < 0.01) (Fig. 3). Although this analysis should be interpreted with caution due to the unbalanced nature of the sample (28 observations between 130 to 265 km and only 6 observations between 1 to 8 km), it suggests that the genetic differentiation among locations only becomes more substantial with increasing distance.

### **DISCUSSION**

In Michigan, producers of tomatoes, peppers, and cucurbits have experienced increasing losses to P. capsici during the last 10 to 15 years. Land suitable for vegetable production is limited in some areas and the length of crop rotation is restricted. A minimum of 3-years rotation to nonsusceptible hosts is a standard recommendation (20). The efficacy of rotation in a disease management program depends on the ability of P. capsici to survive and move among locations. Determining the survival period of naturally produced P. capsici propagules is difficult (2) because inoculum may be present in a small, often undetectable amount. Although significant local spread via water has been demonstrated (21), there is little information concerning the movement of P. capsici among geographically separated locations. We report on isolates from seven geographically separated locations as part of an ongoing investigation aimed at determining how P. capsici survives and characterizing the dynamics of dispersal. Segregation analysis of 17 of the AFLP markers used in this study suggests they are generally inherited as diallelic Mendelian characters and therefore are useful for estimating population genetic measures with P. capsici.

Earlier studies suggest that outcrossing is an important component of the life history of P. capsici and that recombination has a significant impact on the genetic structure of populations (9,10). The data reported here support these previous conclusions, but suggest that outcrossing occurs on a local scale. This is best illustrated by the grouping of isolates into location-specific clusters. Gene flow among locations serves as a powerful evolutionary force to reduce genetic differentiation (6), and the distinct grouping of isolates based on location is typical for populations that are reproductively isolated. It is unlikely that incompatibility among the isolates from different locations is responsible because the progeny from the interregional cross (OP97 × SFF3) were all hybrid and previous crosses between isolates from separate populations suggested similar results (9). The estimated pairwise fixation indices and AMOVA analysis quantified the differences among the populations and indicated that >25% of the observed genetic variation was unique to single locations. Hartl and Clark state that migration of a small number of individuals (e.g., one to two) per

generation is generally sufficient to keep fixation indices at 0.10 or less (6). The observed pairwise fixation indices among the populations presented here suggest that movement among locations was rare. Although polycyclic disease development appears to play an important role in disease development within a single growing season, there were no members of the same clonal lineage recovered among the seven locations or among years at SW1-A or SW1-B.

The finding that movement among locations appears to be rare suggests that the efficacy of rotation may depend more on the long-term survival of P. capsici than on movement among locations. The fields at farm SW1 provided a unique opportunity to investigate survival and spread. Both SW1-A and SW1-B had P. capsici epidemics in 1999, and the only difference among the two was previous rotation patterns. SW1-A was continuously cropped to squash from 1997 to 1999. Location SW1-B was the site of a severe P. capsici epidemic on squash in 1994 that was followed by a soybean and corn rotation until squash was planted again in 1999. The locations are irrigated from separate wells, do not share drainage water, and plant tissue is not knowingly moved among the sites. These fields are of particular interest because they differed significantly in the proportion of mefenoxam insensitive isolates collected in 1999. Only 2 of the 141 genetically unique isolates collected from SW1-A were sensitive to mefenoxam, whereas 24 of the 35 unique isolates recovered from SW1-B were sensitive to mefenoxam. This suggests very little, if any, movement of isolates from SW1-A to SW1-B. The patterns of diversity at the DNA level clearly separate the isolates into two discrete populations and effectively rule out gene flow in 1999. The genic stability of P. capsici at SW1-A from 1998 to 1999, and at SW1-B from 1999 to 2000, suggests that movement into these sites was/ rare. In light of these findings, a reasonable explanation for the epidemic at SW1-B in 1999 is that oospores formed during the 1994 epidemic remained dormant over five winters and provided the initial inoculum. There are reports of oospores surviving extended periods for other Phytophthora spp. (5), and continued tracking of the P. capsici populations at the locations presented here should help us decipher the relative contributions of reintroduction and survival.

The finding that population differentiation increased with distance, considering the magnitude of genetic differentiation at even the closest sites, is consistent with rare founding events originating from nearby locations or from a similar source population. For example, farms SC1 and SC2 are not connected by waterways, nor do they share equipment, but both produce cucumbers for the pickling industry and utilize the same processing station.

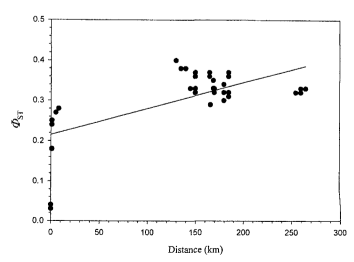


Fig. 3. Association between the degree of genetic subdivision (pairwise  $\Phi_{ST}$ ) and pairwise geographical distances among *Phytophthora capsici* populations at seven locations in Michigan ( $r^2 = 42.6$ ; P < 0.01).

When a semitrailer of cucumber fruit is delivered to the processing station the weight of oversized, undersized, or diseased cucumbers (culls) is estimated and the trailer is reloaded with a corresponding weight of culls sorted from previous deliveries. Traditionally, these culls are spread onto fields with a manure spreader. A single cucumber cull infected with A1 and A2 CT may contain thousands of oospores (K. H. Lamour and M. K. Hausbeck, unpublished data) and it is possible that transfer of infected culls may have contributed to the dissemination of P. capsici in Michigan. All of the locations sampled in this study had a history of P. capsici epidemics and investigation of a newly infested field should provide insight into the nature of founding events.

In summary, it appears that *P. capsici* persists in Michigan fields as reproductively isolated outcrossing populations in which the sexual stage is effectively linked to long-term survival. Thus, even though single genotypes have the potential to increase significantly within a single season, genic diversity is maintained over time and new gene combinations are constantly generated. Comparison of future sample sets to the baseline data presented here should provide an opportunity to further clarify the contributions of movement among locations and survival to the population structure of *P. capsici* in Michigan.

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## Worksheet 3-A(9)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Crop Rotation		y: The spatiotemporal genetic structure of	
	-	Phytophthora capsici in Michigan and implications	
		for disease management.	

## Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the use of this alternative on your site 2

miere any resultant openine restrictions that minibit the	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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## Worksheet 3-A(9)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. K.H. Lamour 2. Author(s) or researcher(s) M.K. Hausbeck Phytopathology 92:681-684, 2002 3. Publication and Date of Publication Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Crop Rotation 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Crop rotation is not highly effective because both mating types of *Phytophthora capsici* are present in Michigan fields, resulting in an oospore capable of surviving for long periods of time. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable since the research was conducted in Michigan, USA.

OMB Control # 2060-0482

## New Frontiers in Plant Disease Losses and Disease Management

# The Spatiotemporal Genetic Structure of *Phytophthora capsici* in Michigan and Implications for Disease Management

K. H. Lamour and M. K. Hausbeck

Department of Plant Pathology, Michigan State University, East Lansing 48824. Accepted for publication 12 February 2002.

Root, crown, and fruit rot caused by *Phytophthora capsici* Leonian is a limiting factor for the production of peppers, tomatoes, and cucurbit crops in Michigan and the United States. Like many species in the genus *Phytophthora*, *P. capsici* has the potential for rapid polycyclic disease development from a limited amount of initial inoculum (6). *P. capsici* produces caducous sporangia that can be spread by wind-blown rain or release 20 to 40 motile zoospores in the presence of free water. The polycyclic phase of disease development is thought to be driven primarily by asexual spore dispersal at a local scale (within and down rows). Sexual reproduction requires both the A1 and A2 compatibility types (CTs) and results in the production of thick-walled oospores. Oospores are thought to serve as the primary survival structure outside of host tissue.

Recommended disease management strategies stress the importance of avoiding excess water in the plant rhizosphere by using well-drained fields, conservative irrigation, and planting on raised beds. Additional recommendations include rotation to nonsusceptible hosts for at least 2 years and the use of fungicides. The phenylamide fungicide (PAF) mefenoxam is a systemic compound with high activity against P. capsici and has been used by growers throughout the United States to control P. capsici. Insensitivity to PAF has been reported for a number of other oomycetous organisms (Bremia lactucae, P. infestans, and P. sojae, etc.) and appears to be conferred by a single incompletely dominant gene of major effect (1). Growers in Michigan practicing 2+-year rotation in well-drained fields using an array of fungicidal management tools have experienced significant losses to P. capsici. Michigan is the number one producer of cucumbers for pickling in the United States and it was at the request of grower groups associated with this industry that research into the epidemiology and reproductive biology of *P. capsici* on cucurbit hosts was initiated.

Although many researchers cite oospores as the most likely propagule for survival outside of host tissue, there have been very few investigations specifically aimed at determining the impact of sexual reproduction in natural populations of *P. capsici*. Our hypothesis was that the sexual stage may play an important role not only in survival but also in the adaptation of *P. capsici* populations to environmental stresses (e.g., fungicides). Our goal was to perform a comprehensive investigation of the phenotypic and genetic diversity present in *P. capsici* populations from the major vegetable production regions of Michigan, with the implicit intention of addressing questions concerning epidemiology, repro-

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ductive biology, and the durability of currently recommended management strategies.

#### **METHODOLOGY**

Isolate collection and maintenance. Sampling of diseased fields began at the end of the 1997 growing season and continued through September 2000. In all cases, fields were sampled on a grid with quadrants varying from 40 m² to 12 km². A limited number of isolates were collected in 1997. In 1998, the strategy was to collect as many samples from as many fields as possible. This strategy was modified in 1999 and 2000 to focus on specific fields. Isolations from diseased plants were made onto selective media and single zoospore cultures were generated according to standard single sporing techniques (3). Isolates were placed into long-term storage (15°C) using a hemp seed/sterile water technique.

Phenotypic characterization. Single zoospore isolates were screened for CT using known A1 and A2 isolates. In vitro screening techniques published for other Phytophthora species for assessing sensitivity to mefenoxam were compared and a novel, simple, high dose screen using 100 ppm of mefenoxam-amended V8 agar was found to separate field isolates into three modal distributions that appeared consistent with the expectations of a single incompletely dominant gene governing mefenoxam insensitivity (e.g., sensitive, intermediately sensitive, and fully insensitive). These putative mefenoxam sensitivity (MS) groupings were tested by performing a series of crosses and testing whether the observed progeny sets met the expectations for Mendelian inheritance of a single incompletely dominant gene controlling insensitivity to mefenoxam. Sexual crosses were conducted on unclarified V8 agar plates and incubated for 3 months in the dark. Individual germinated oospores were recovered after 3 months using previously published techniques (2).

The efficacy of this in vitro mefenoxam screening technique was further tested in pumpkin seedlings using progeny from a cross between parents intermediately sensitive to mefenoxam. Nine isolates from each of the three MS categories were screened for pathogenicity on untreated seedlings. Single sensitive, intermediately sensitive, and fully insensitive isolates were then placed onto the unwounded surface of plants treated with either a field rate of mefenoxam, three times the field rate, or distilled water. Lesion diameters on seedling stems were measured after 4 days.

Genetic characterization. Single zoospore isolates were grown in antibiotic-amended V8 broth for 3 days at room temperature. Mycelial mats were washed, frozen, lyophilized, and ground with a sterile mortar and pestle. DNA was extracted with either a Qiagen Dneasy extraction kit (Qiagen, Valencia, CA) or via a cetyltrimethylammonium bromide (CTAB) procedure. A variety

of methods for generating molecular markers were tested for efficacy including isozyme, random amplified polymorphic DNA, and amplified fragment length polymorphism (AFLP). The AFLP technique resulted in a large number of reproducible markers and was chosen to characterize samples of P. capsici from Michigan. The AFLP technique involves cutting genomic DNA with moderately rare cutting (EcoRI) and frequent cutting (MseI) restriction enzymes, while concomitantly ligating synthetic adaptor fragments of DNA to the sticky ends created by the restriction enzymes (7). The result is a large number of DNA fragments that have ends with known DNA sequences. Amplification of fragment subsets (termed fingerprints) can be accomplished using polymerase chain reaction (PCR) primers complementary to the adaptor sequences with additional "selective" nucleotides. Changing the amount and type of selective nucleotides results in different subsets or fingerprints. Stringent PCR cycling parameters (touchdown technique) are used to ensure the fidelity of the reaction. For the analysis summarized here, adaptor sequences and fluorescent labeled selective primers were purchased as a kit through Perkin-Elmer ABI (Applied Biosystems, Foster City, CA). Using this system, AFLP fragments were resolved on a polyacrylamide gel by an ABI 377 gene sequencer. Fluorescent labels were excited by a laser and band emissions were analyzed in the form of an electropherogram where peaks represent individual bands. The sizing of fragments was particularly robust because a DNA ladder was loaded with every sample into the gel. To test for the reproducibility of fingerprints, DNA was extracted from a single isolate on three separate occasions approximately 3 months apart and subjected to the aforementioned protocol.

Data analysis. Isolates with identical multilocus AFLP fingerprints were considered to be members of the same clonal lineage and only a single representative was used for analysis. Because AFLP markers can only be scored confidently for presence (1) or absence (0), allele frequencies were estimated based on the assumption that populations under investigation meet the criterion for Hardy-Weinberg equilibrium, and that loci have only one "present" allele. The term population refers to all samples taken from a single field during a single year.

Genetic diversity within single populations was assessed by calculating the average number of polymorphic bands and estimating the average heterozygosity. Fixation indices were calculated according to methods of Weir and Cockerham (8) for populations from the same site over multiple years and among populations in Michigan using the program tools for population genetic analysis (TFPGA) (M. P. Miller, Northern Arizona University, Flagstaff). Confidence intervals for F statistics at the 95% confidence level were generated by bootstrapping at 1,000 iterations. The program NTSYS-pc version 2.02k (Exeter Software, Setauket, NY) was used to construct a similarity matrix from the presence/absence (1/0) data. Cluster analysis using the unweighted pair group with arithmetic averages (UPGMA) method was performed on the matrix and a tree was generated to give a visual representation of isolate similarity. Excoffier's ARLEQUIN program (L. Excoffier, University of Geneva) was used to assess population differentiation using a phenetic approach termed analysis of molecular variance (AMOVA), which allows for total genetic variation to be partitioned within and among populations using a classical analysis of variance (ANOVA).

## **RESULTS**

Phenotypic results. Five isolates were recovered in 1997 from five different farms (four A1 and one A2 CT). One isolate was fully insensitive to mefenoxam, whereas the other four were fully sensitive. These findings prompted the extensive sampling conducted in 1998 in which 523 isolates (473 from cucurbits and 30 from bell pepper) were collected from 14 farms. A frequency histogram plotting percent growth of control on 100 ppm of

mefenoxam-amended media versus number of isolates revealed a trimodal distribution (3). Putative MS categories were assigned based on these groupings with sensitive (S) <30% growth of control, intermediately sensitive (IS) between 30 and 90% growth of control, and insensitive (I) >90% growth of control. In vitro crosses between isolates representative of the different putative sensitivity categories (S  $\times$  S, I  $\times$  S, IS  $\times$  S, and IS  $\times$  IS) resulted in progeny sets not significantly different than expected for insensitivity inherited as a single incompletely dominant gene unlinked to CT (P = 0.05) (3). In 1998, 55% of the isolates were sensitive to mefenoxam, 32% were intermediately sensitive, and 13% were fully insensitive to mefenoxam. A1 and A2 CTs were recovered in a ratio of approximately 1:1 in 8 of the 14 farms. Oospores were detected in naturally diseased cucurbit fruit from four farms, and 223 oospore progeny were recovered and germinated from a single diseased cucumber. All six possible MS × CT combinations were detected in this naturally occurring oospore progeny set (3).

In planta studies using sensitive, intermediately sensitive, and fully insensitive *P. capsici* isolates supported the in vitro screening categories, with sensitive isolates causing no disease on mefenoxam-treated plants, intermediately sensitive isolates being slowed by mefenoxam, and fully insensitive isolates showing no difference in the ability to colonize host tissue between treated and untreated plants at three times the field rate. All the progeny isolates were pathogenic on untreated pumpkin plants (K. H. Lamour and M. K. Hausbeck, *unpublished data*).

Sixty-three mefenoxam insensitive (18% intermediate and 82% fully insensitive) isolates were recovered from a single southwest Michigan field in 1998. Field experiments were conducted in this field during 1999 and 2000, testing alternative cultural control strategies, and no mefenoxam was applied. Two hundred isolates were recovered from this site over the course of the 1999 season and 34 isolates at the beginning of the 2000 season. Of the 200 isolates recovered in 1999 from this field, 141 had unique AFLP genotypes. Seventy percent of these were fully insensitive to mefenoxam, 28% were intermediately sensitive, and 2% were sensitive. In 2000, 15% of the isolates were intermediately sensitive and 85% were fully insensitive. A single fully insensitive clonal lineage rose in frequency over the course of the 1999 season and comprised 20% of the total number of samples recovered (4).

During 1999 and 2000, approximately 2,500 isolates were recovered from farms in Michigan. Both the A1 and A2 CTs were present in every field sampled, and mefenoxam insensitivity was detected in the majority of farms that had a history of mefenoxam use.

Genetic results. Nine populations from the four major vegetable production areas of Michigan were analyzed with the AFLP procedure (N = 641). AFLP analysis resolved a total of 94 clearly discernable markers when considering all the isolates together. No single isolate or group of isolates from a single location contained all 94 markers. The total number of AFLP loci in a single population ranged from 68 to 80. Seventeen (18%) fragments were fixed for the present state across all populations, 12 (13%) fragments were polymorphic in all populations, and 65 (69%) were fixed for presence or absence in some populations and polymorphic in others. The number of polymorphic bands within a single population ranged from 37 to 46 with estimated heterozygosities ranging from 0.18 to 0.22. Clonal reproduction was significant within single fields over the course of the growing season. For example, genotypic diversity in a single field ranged from 100% at the beginning of the growing season (seedling stage) to <30% at the time cucurbit fruit were ready for harvest (4). When considering all nine populations, genotypic diversity ranged from 42 to 96% with an average of 74% of the isolates in any sample set having unique genotypes. Although clonal reproduction was significant within single fields within years, no clones were recovered from single fields between years or among fields separated by at least 1 km. Fixation indices  $(\phi_{ST})$  between the

populations sampled on consecutive years were very close to zero, indicating that gene diversity was not measurably impacted by genetic drift (5). The overall estimated  $\phi_{ST}$  for populations from different locations was 0.35, indicating that approximately 35% of the total genetic diversity present in Michigan *P. capsici* populations is found among populations and 65% is found within any one population. AMOVA partitioned genetic diversity among (40%) and within (60%) populations. The similarity tree based on UPGMA cluster analysis clearly showed that isolates from the

same site sampled over years branched from the same node, with no clustering of isolates based on the year of sampling. Cluster analysis also clearly showed that populations separated geographically branched from population-specific nodes (5).

## **DISCUSSION**

During the past 10 years, Michigan has experienced a steady increase in the incidence of root, fruit, and crown rot on cucurbits

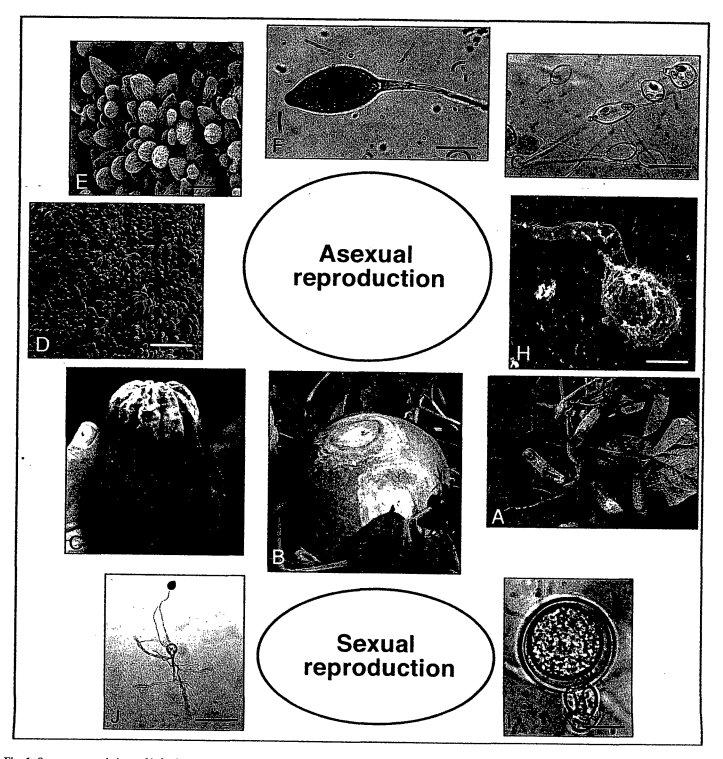


Fig. 1. Spore types and signs of infection caused by *Phytophthora capsici* on cucurbit fruit: A, infected cucumber, B, pumpkin, and C, acorn squash fruit. D, Scanning electron microscope (SEM) photo of an infected cucumber showing tufts of sporangia produced on the surface of the fruit (Bar = 300  $\mu$ m). E, Close-up of a single tuft of sporangia (Bar = 30  $\mu$ m). F, Typical papillate sporangium with a long pedicel (Bar = 20  $\mu$ m). G, Zoospores exiting sporangia after immersion in water (Bar = 50  $\mu$ m). H, SEM photo of a single encysted zoospore that germinated and directly penetrated the epidermis of a cucumber fruit (Bar = 4  $\mu$ m). I, Typical amphigynous oospore (Bar = 10  $\mu$ m). J, A germinating oospore with multiple germ tubes and a terminal sporangium (Bar = 100  $\mu$ m).

caused by *P. capsici*. Rotation to nonsusceptible hosts, in conjunction with cultural and chemical control strategies, have not provided economic control. Correspondence with other vegetable pathologists suggests that this phenomenon is not confined to Michigan, and a similar increase in control failures due to blight by *P. capsici* is being reported throughout the United States.

Investigation of the inheritance of MS demonstrated that MS is inherited as a single incompletely dominant gene unlinked to CT. In 1998, all six possible MS × CT combinations were present in single fields and insensitivity to mefenoxam was common in Michigan. Typical amphigynous oospores were observed in *P. capsici*-infected cucurbit fruit from multiple locations, and oospore progeny from a single naturally infected fruit showed segregation for MS and CT. These findings strongly support the hypothesis that sexual reproduction is occurring in the field, and also suggest that sexual recombination may directly generate progeny fully insensitive to mefenoxam. Tracking a single mefenoxam insensitive population over 2 years in the absence of mefenoxam selection pressure suggests that costs associated with mefenoxam insensitivity are minimal.

Estimates of average heterozygosity and polymorphism indicate surprisingly high levels of gene and genotypic diversity in all the populations of P. capsici analyzed. Tracking-a single population through an entire growing season showed that asexual reproduction plays a significant role in disease development within a single season. Sampling single fields over consecutive years suggested that clones do not survive Michigan winters and that oospores are the primary survival propagule. Estimation of fixation indices for samples from the same site over consecutive years suggested that there was not a significant reduction in genetic diversity between growing seasons. This implies that populations are large enough to withstand dramatic effects of genetic drift. Cluster analysis revealed unambiguous groups corresponding to geographical locations with regional populations showing more similarity overall than populations from different regions. Population pairwise fixation indices corroborated this finding. The estimated overall fixation index and AMOVA are in agreement with both, suggesting that most (approx 60%) of the total genetic variability in Michigan is found within any one population, but that a relatively large component (40%) of genetic variability is found among populations.

Recommendations based on our findings are as follows: (i) the fungicide mefenoxam may be of limited usefulness because insensitivity appears to be selected for rapidly and is unlikely to decrease when mefenoxam selection pressure is removed; (ii) fields with epidemics are likely to harbor oospores for an extended amount of time (at least 5 years), and this factor must be considered before replanting to susceptible hosts; and (iii) factors that may contribute to the introduction of *P. capsici* into uninfested fields (e.g., drainage ditches between farms, irrigation ponds, and the dumping of culls) need to be considered and if possible avoided, because once an epidemic is established we have found no evidence that the population will become extinct in an agriculturally meaningful time period.

From an evolutionary perspective, it is clear that *P. capsici* has successfully colonized a number of geographical locations in

Michigan and that each of the populations sampled thus far have similarly high levels of genetic variability. The genetic stability of single populations over multiple years, the high fixation indices between even geographically close populations (1 km), and the clear structuring based on UPGMA cluster analysis all suggest that long-distance dispersal of inoculum is not common and that geographically isolated populations are also genetically isolated. It appears that the sexual stage of the P. capsici life cycle plays a significant role in survival as well as maintaining both genic and genotypic diversity, and has likely played a key role in the evolution of mefenoxam insensitivity. The combination of high levels of genetic variability, thick-walled oospores, and polycyclic asexual disease development make P. capsici a formidable pathogen (Fig. 1). This work underscores the need for management strategies aimed at preventing the spread of P. capsici to uninfested field sites and suggests that management strategies aimed at limiting spread within a single season may be the only option for growers with P. capsici-infested fields.

## **ACKNOWLEDGMENTS**

This work was funded by the Michigan Agricultural Experiment Station, Michigan State University Extension, Michigan Department of Agriculture, Michigan Farm Bureau (GREEN cooperative), Pickle and Pepper Research Committee, Pickle Packers International, Inc., and the Pickle Seed Research Fund, Pickle Packers International. We thank M. Bour, C. Hunter, J. Jabara, P. Tumbalam, E. Webster, and J. Woodworth for competent laboratory assistance. K. Lamour thanks his Ph.D. committee members A. Jarosz, R. Hammerschmidt, and F. Trail for guidance and extends sincere thanks to M. Hausbeck for fulfilling her role as mentor in an exemplary manner.

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Worksheet 3-A(9)(e). Alternatives - Techni	ical Feasibility of Alternatives to Methyl Bromide
	ive pest management strategy on the list (see previous page) is or is ains 9 questions. You must complete one copy of worksheet 3-A for by bromide alternative. Use additional pages as need.
the worksheets as follows. For the same alternative, fi alternative, second research study, label the workshee	for each alternative, for each research study addressed. Please number irst research study, label the worksheet 3-A(1)(a). For the same at 3-A(1)(b). For the first alternative, third research study, label the esearch study, label the worksheet 3-(A)(2)(a). For the second at 3-(A)(2)(b).
When completing Section II, if you cite a study that is	on the EPA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the	e Research Summary Worksheet.
	ed in this worksheet in a narrative review of one or more relevant ection I and questions 1 through 8 in Section II. A Research Summary for each study reviewed.
BACKGROUND	
successfully instead of methyl bromide by crop and circum	ares (pesticide and non-pesticidal, and their combination) could be used estance (geographic area.) The Agency has developed a list of possible can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.
There are three major ways you can provide the Agency w (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website	ith proof of your investigative work.
scientifically sound manner. The studies should include a c	developed by others, it is important that the studies be conducted in a description of the experimental methodology used, such as application rates, trieties of the crop used, etc. All results should be included, regardless of A unless they are listed on the Agency website.
	y of crops on its website and knows of more studies currently in progress. y available. You are encouraged to review the EPA website and other nic area.
· · · · · · · · · · · · · · · · · · ·	ces, some alternatives are not technically feasible and therefore no research in Seattle). You should look at the list of alternatives provided by the Agency your geographic area.
Use addi	tional pages as needed.
Alternative: Crop Rotation	Study: Investigating the impact of crop rotation on the genetic structure of <i>Phytophthora capsici</i> .
Section I. Initial Screening on Technic	cal Feasibility of Alternatives
	in the second of
Are there any location-specific restrictions that inhib	·
1a. Full use permitted 1b. Township caps	X
is. Township caps	

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

1c. Alternative not acceptable in consuming country

1d. Other (Please describe)

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## Worksheet 3-A(9)(e). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. K.H. Lamour 2. Author(s) or researcher(s) M.K. Hausbeck submitted for publication in Fungal Genetics and Biology, 2002 3. Publication and Date of Publication Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Crop Rotation, Fallow 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Crop rotation was not effective because the oospore is long lived in Michigan soils. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable since the research was conducted in Michigan, USA.

OMB Control # 2060-0482

## K. H. Lamour, Page 1, Fungal Genetics and Biology

# Investigating the impact of crop rotation on the genetic structure of *Phytophthora capsici*

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1 ABSTRACT

Lamour, K. H. and Hausbeck, M. K. 200-. Investigating the impact of crop rotation on the genetic structure of *Phytophthora capsici*. Fungal Genetics and Biology : -

Phytophthora capsici isolates (N = 104) recovered from a single field planted to cucumbers (1998), corn (1999 and 2000), and tomatoes (2001) were screened for compatibility type, mefenoxam sensitivity, and AFLP profiles. Eighty-nine percent of the isolates had unique genetic profiles with 60% of the AFLP markers polymorphic. The ratio of A1:A2 compatibility types was  $\approx 1:1$  and the frequency of mefenoxam insensitive isolates was similar between years. No clonal lineages survived between 1998 and 2001 and the pool of phenotypic and genetic diversity remained essentially intact. This suggests that (i) the site harbors a discrete outcrossing population with little immigration, (ii) two years rotation to corn did not significantly reduce the effective population size, and (iii) cropping to tomatoes did not significantly impact the overall genetic structure of P. capsici at this location. The importance of sex in maintaining diversity and allowing survival within a natural population of P. capsici is discussed.

**Index descriptors:** sex, recombination, genetic drift, host selection, migration, AFLP, population genetics, temporal variation.

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## INTRODUCTION

Phytophthora capsici has been responsible for significant losses to vegetable producers in the United States during the last century (1, 7-9, 13, 19, 21, 22). In Michigan, P. capsici causes root, crown, and fruit rot on cucumbers, squash, pumpkins, tomatoes, and peppers and the incidence and severity of disease have increased significantly in the last ten to fifteen years. Control strategies employed by Michigan vegetable producers include planting at well-drained sites, crop rotation to non-susceptible hosts for at least two years, and the application of fungicides. None of the above control strategies have provided economic control under optimal environmental conditions for disease development. The phenylamide fungicide (PAF) mefenoxam, or the similarly acting compound metalaxyl, has been used by some growers. Although PAFs are fungistatic to sensitive isolates of P. capsici, a significant problem is that many populations of P. capsici in Michigan and elsewhere exposed to metalaxyl or mefenoxam have adapted to PAF selection pressure and the efficacy of PAF's in these populations may be greatly reduced (2, 3, 10, 11, 17, 18).

Phytophthora capsici is an outcrossing diploid organism that requires the presence of both A1 and A2 compatibility types to complete the sexual stage and produce oospores (4). Oospores are thick-walled and appear to play an important role in survival. Asexual reproduction is completed by a coenocytic mycelial thallus able to produce sporangia on the surface of infected tissue. Sporangia are borne on long caducous pedicels and may germinate directly or indirectly. Sporangia germinate indirectly when immersed in free water and produce 20 to 40 motile zoospores. As is the case with many members of the genus Phytophthora, excess moisture favors epidemic development (20).

Reports on the spatiotemporal genetic structure of P. capsici in Michigan suggest that epidemics are initiated by dormant genetically diverse inoculum and that movement between geographically separated locations is not common (12). Clonal reproduction may be significant within a single growing season, but it appears that the perpetuation of clonal lineages is limited to single fields and single growing seasons (12).

Previous temporal studies were conducted on Michigan P. capsici populations infecting cucurbit hosts over the course of a single growing season or among years separated by a single

winter (eg; November to March) and may not reflect the dynamics of *P. capsici* over longer periods of time (eg; a typical two year rotation) or among diverse hosts. In the present investigation we test the hypothesis that *P. capsici* can survive a thirty month non-host period via dormant genetically diverse propagules. In addition, we investigated the effects of selection among cucurbit and solanaceous hosts, genetic drift, and migration on population structure.

## MATERIALS AND METHODS

**Isolate recovery:** Tomato plants and cucumber fruit exhibiting typical signs and symptoms of infection by *P. capsici* were collected from a vegetable production field located in south central Michigan that was planted to pickling cucumbers in 1998, corn in 1999 and 2000, and processing tomatoes in 2001. Both the cucumbers and tomatoes were planted on flat ground and the field was irrigated via a pivot irrigation system supplied with water from a well. At the time of harvest in 1998, the majority of the cucumber fruit present in this field had obvious signs and symptoms of disease ranging from discrete water-soaked lesions to being entirely covered with a white, powdery, layer of sporangia. Other than the fruit, the plants appeared to be healthy with no symptoms of disease on foliage, vines, or stems. Close inspection indicated that a limited number of plants (< 5%) scattered throughout the field were stunted.

In 2001, the most frequent above-ground symptom on tomatoes infected with *P. capsici* was stunting with a small number of the infected plants showing wilt symptoms. Foliar lesions were not observed. Plants were recovered prior to fruit being set and the incidence of fruit infection was not determined. The crown area of infected plants was brown-black. Infected plants often had a brown crumbly epidermis from the soil line to the tap root and a significant reduction in feeder roots. In some cases, plants with infected tap roots had numerous adventitious roots above the point of infection. During both 1998 and 2001 diseased plant material was collected in haphazard fashion throughout the field.

Infected cucumbers were snapped in half by hand and a small section (c. 1 cm<sup>2</sup>) of tissue removed from beneath the cuticle. The root and crown area of infected tomato plants were rinsed with tap water and patted dry with paper towels before a section of tissue at the edge of an expanding lesion was removed. Tissue was not surface sterilized prior to isolation. Isolations were made onto BARP (benomyl 25 ppm, ampicillin 100ppm, rifampicin 30ppm, and

pentachloronitrobenzene 100ppm) amended UCV8 (840 ml distilled water, 163 ml unclarified V8 juice, 3 g CaCO<sub>3</sub>, and 16 g agar) plates. Plates were incubated at room temperature in the dark for 2 to 3 days before colonies were transferred. Procedures for obtaining single zoospore isolates were as previously described (10). Single zoospore cultures were maintained on RA (rifampicin 30 ppm, ampicillin 100 ppm)-UCV8 plates and transferred bi-monthly. For longterm storage, a 7-mm plug of expanding mycelium from each culture was placed into a 1.5 ml microfuge tube with one sterilized hemp seed and 1 ml of sterile distilled water (SDW). Isolates were then incubated for 2 to 3 weeks at 23 to 25°C before being stored at 15°C. Compatibility type and mefenoxam sensitivity determination: Agar plugs from the edge of an expanding single-zoospore colony were placed at the center of UCV8 plates approximately 2 cm from field isolates OP97 (A1) and SP98 (A2) and incubated in the dark at 23 to 25°C for 3 to 6 days. Following incubation, compatibility type was determined. Agar plugs from the edge of actively expanding single-zoospore colonies were placed at the center of 100 x 15 mm UCV8 plates amended with 0 or 100 ppm mefenoxam (Ridomil Gold EC, Novartis, Greensboro, NC; 48% AI, suspended in SDW; added to UCV8 cooled to 49°C). Inoculated plates were incubated at 23 to 25°C for 3 days and colony diameters measured. Percentage growth of an isolate on amended media was calculated by subtracting the inoculation plug diameter (7-mm) from the diameter of each colony and dividing the average diameter of the colony on amended plates by the average diameter of the colony on unamended control plates. All tests were conducted at least twice. An isolate was scored as sensitive (S) if growth at 100 ppm was < 30% of the control, intermediately sensitive (IS) if growth was between 30 and 90% of the control, and insensitive (I) if growth was > 90% of the control (10). DNA extraction and AFLP fingerprinting: Bacterial contamination was avoided by using a modified Van Teigham cell (4). The uppermost portion of a 7-mm plug of mycelium was placed onto the surface of RA-WA plates (rifampicin 30 ppm, ampicillin 100 ppm, 1000 ml distilled water, and 16 g agar) and an autoclaved cap from a 1.5 ml microfuge tube was placed over the plug which forced the isolate to grow through the amended medium. Isolates were incubated in the dark for 2 to 3 days before two 7-mm plugs were transferred to approximately 15 ml of

RA-UCV8 broth in 100 x 15 mm Petri dishes and incubated in the dark for three days at 23 to

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25°C. Mycelial mats were washed with distilled water and dried briefly under vacuum before being frozen to -20°C and lyophilized.

Lyophilized mats were ground with a sterile mortar and pestle. Whole genomic DNA from approximately 50 mg of ground mycelium was extracted using a QIAGEN Dneasy Plant Mini Kit (QIAGEN Inc., Valencia, CA) according to the manufacturers directions. DNA was quantified using Nucleic Acid QuickSticks (CLONTECH, Palo Alto, CA) according to the manufacturer's directions or on 1.5% agarose gels. Approximately 100 ng of DNA was then subjected to a restriction/ligation reaction, pre-selective amplification, and selective amplifications using the PCR core mix, adaptor sequences, core primer sequences and fluorescence labeled primers provided in the AFLP<sup>TM</sup> Microbial Fingerprinting Kit (Perkin-Elmer Corp., Foster City, CA) and performed exactly as described in the PE/ABI AFLP Microbial Fingerprinting protocol part # 402977 Rev A (23). All PCR reactions were performed using an MJ Research Minicycler (MJ Research Inc., Waltham, MA) in 0.2 ml tubes according to the cycling parameters outlined in the microbial fingerprinting protocol.

An initial optimization set of reactions was performed using pre-selective products from *P. capsici* isolate OP97 which was isolated from a cucumber fruit in 1997 (10). Amplifications with the selective primers EcoRI-AA, AC, AG and AT were performed in all 16 combinations with the MseI-CA, CC, CG and CT selective primers. EcoRI selective primers were labeled at the 5' end with either carboxyfluorescein (FAM), carboxytetramethyrhodamine (TAMRA), or carboxy-4',5'-dichloro-2',7'-dimethoxyfluorescein (JOE) fluorescent dyes. The fluorescent dyes were excited by laser radiation and visualized by their characteristic absorption-emission frequencies. Only the fragments containing an EcoRI restriction site were resolved.

Selective amplification AFLP products and a carboxy-X-rhodamine (ROX) size standard were loaded into each lane on a denaturing polyacrylamide gel and the fragments resolved in an ABI Prism 377 DNA Sequencer. Results were prepared for analysis in the form of electropherograms using GeneScan Analysis software (PE/ABI). AFLP fragments were scored manually as present = 1 or absent = 0 using Genotyper (PE/ABI). Only DNA bands which consistently exhibited unambiguous presence/absence profiles were scored.

In order to assess the reproducibility of AFLP profiles, a single isolate, OP97, was

subjected to the aforementioned protocol using three optimal primer pair combinations on three separate occasions approximately three months apart.

Clone detection: AFLP fragments for each field isolate were scored for presence or absence, and the binary data matrix was converted to a similarity matrix with the program NTSYSpc version 2.02k (Exeter Software, Setauket, NY). Unweighted pair group method with arithmetic averages (UPGMA) cluster analysis was performed on the similarity matrix and a tree was generated. Isolates showing complete homology at all loci were considered to be members of the same clonal lineage, and except for a single representative isolate (referred to as a clone) were excluded from population genetic analysis (15).

Population genetic analysis: Sample sets collected from single fields during a single year were considered a population. Populations were assumed to be in Hardy-Weinberg equilibrium, and each AFLP locus was assumed to be di-allelic and selectively neutral. The program 'Tools for population genetic analysis' (TFPGA) (Miller, M. P., Northern Arizona Univ., Flagstaff, AZ) was used to assess genetic diversity within each population on the basis of estimated average heterozygosity (16) and the proportion of polymorphic loci at the 95% level (6), and to calculate pairwise and overall fixation indices (F-statistics) according to the methods of Weir and Cockerham (24). Confidence intervals for F-statistics at the 95% confidence level were generated by boot-strapping using 1000 iterations. Estimates of the percent polymorphic loci and estimated average heterozygosity were calculated based on the 68 AFLP markers resolved.

The fixation index, as described by Wright, equals the reduction in heterozygosity expected with random mating at any one level of a population hierarchy relative to another, more inclusive level of the hierarchy (25). Weir and Cockerham's approach to estimating fixation indices attempts to correct for the effects of sampling a limited number of organisms from a limited number of populations and is reported as  $\Phi_{ST}$  instead of  $F_{ST}$  (24). Theoretically, the fixation index has a minimum of 0 (no loss of heterozygosity between the populations compared) and maximum of 1 (indicating fixation for alternative alleles in different populations or a total loss of heterozygosity), but, as discussed by Hartl and Clark (6), the observed maximum is usually much less than 1. Wright has suggested the following qualitative guidelines for the interpretation of fixation indices: the range 0 to 0.05 indicates little genetic differentiation, 0.05

to 0.15 indicates moderate genetic differentiation, 0.15 to 0.25 indicates great genetic differentiation, and values above 0.25 indicate very great genetic differentiation.

Using the program NTSYS-pc, the combined 0/1 data matrix for isolates from all populations was used to construct a genetic similarity matrix of all possible pairwise comparisons of individuals within and among populations using Jaccard's similarity coefficient: GS(ij) = a/(a+b+c). GS(ij) is the measure of genetic similarity between individuals i and j, where a is the number of polymorphic bands shared by i and j, b is the number of bands present in i and absent in j, and c is the number of bands present in j but absent in i. A tree was constructed using  $\overline{UPGMA}$  cluster analysis to provide a graphic representation of the relationships among isolates.

Genetic structure was also examined by analysis of molecular variance (AMOVA) using the ARLEQUIN software package (Excoffier, L., University of Geneva, Geneva). The AMOVA analysis was used to partition the variance in banding patterns within and among the populations. Significance values were assigned to variance components on the basis of a set of null distributions generated by a permutation process which randomly assigned individuals to populations and drew 1000 independent samples.

RESULTS

RESULIS

**Isolate recovery:** In 1998, 141 isolates of *P. capsici* were recovered from infected cucumber fruit. Phenotypic characterization of all 141 isolates and genetic characterization of 57 isolates has been reported previously (10, 12). Here we report only on the 57 isolates characterized genetically. In 2001, 47 isolates of *P. capsici* were recovered from infected tomato plants. **Genetic diversity, compatibility type, and sensitivity to mefenoxam:** Evaluation of the 16

EcoRI + 2/MseI + 2 selective primer pair combinations indicated that EcoRI + AC/ MseI + CA gave the most clearly resolved fragment profile and was used to amplify genomic DNA from all isolates in both the 1998 and 2001 sample sets. This primer combination resulted in 68 clearly resolved fragments of which 42 fragments were polymorphic in 1998 and 45 were polymorphic in 2001 (Table 1). All 68 fragments were present in both 1998 and 2001 and no novel fragments were detected among years. AFLP profiles for isolate OP97, generated from separate DNA extractions on three separate occasions over a one year period, resulted in identical banding

patterns with the only difference being minor changes in the intensity of the electropherogram signal. Occasionally individual reactions resulted in poorly resolved fingerprint profiles (eg, low intensity of signal) and were repeated until signals were deemed optimal.

In 1998, there were five clonal lineages detected with three lineages comprised of two members and two lineages with three isolates each. In 2001, there were four clonal lineages detected with two members each. No members of the same clonal lineage were detected among the isolates collected in 1998 and 2001. The ratio of A1:A2 isolates was 29:21 in 1998 and 21:22 in 2001 (Table 2). These numbers approximate the 1:1 ratio expected for a randomly outcrossing diploid organism. The percentage of isolates falling into the three mefenoxam sensitivity categories was 40% and 44% sensitive, 56% and 38% intermediately sensitive, and 4% and 18% fully insensitive for 1998 and 2001 respectively (Table 2).

**Temporal dynamics:** The fixation index ( $\Phi_{ST}$ ) among the populations of *P. capsici* recovered in 1998 and 2001 was 0.05 with a standard deviation of 0.01. This indicates that very little genetic differentiation, or loss of heterozygosity, occurred between years at this location. The number and identity of bands polymorphic at the 95% level and the estimated average heterozygosity (0.17 for both years) remained relatively similar over time (Table 1). AMOVA analysis partitioned 3% of the total variability among years indicating that 97% of the variation found in 1998 was also found in 2001 (Table 3). These data suggest that there was enough out-crossing and survival of the resulting recombinant progeny at this location to maintain genic diversity. UPGMA analysis showed that unique genotypes were between 76 and 96% similar and that isolates from 1998 and 2001 were dispersed randomly throughout the tree (Fig 1). There was no grouping of isolates based on year or host.

## Discussion

In this study we investigated the genetic structure *P. capsici* infecting cucumbers and tomatoes separated by two years of crop rotation to corn. Previous studies indicate that genetic diversity is high in natural populations of *P. capsici* in Michigan and that the pool of AFLP markers resolved from the isolates at this location is unique (12). By tracking the changes in the identity and frequency of the AFLP markers over time we were able to gain novel insight into the survivability of *P. capsici* at a naturally infested site and to begin deciphering the impact of crop

rotation on P. capsici's population structure.

There are three major evolutionary forces that could have changed the genetic structure of *P. capsici* at this location between 1998 and 2001; genetic drift, selection, and migration (5, 14). A significant pressure by any one of these should be discernable in the patterns of genetic diversity recovered among years. Genetic drift refers to the random sampling process that occurs within small populations over time (6). If only a small number of *P. capsici* propagules survived between the epidemics in 1998 and 2001, then there should be significant changes in the respective frequencies of neutral genetic markers just due to chance. In particular, it's expected that the total genetic diversity recovered in 2001 would be a subset of that recovered in 1998 because some rare markers would be missed in the sampling (= survival) process. An in-depth summary of genetic drift is not possible in this context, but it is clear that there was not a significant reduction in the total genetic diversity among years at this location. This is illustrated by the fixation index estimate of 0.05 and the AMOVA analysis which indicate that between 95 and 97% of the genetic diversity found in one year was also found in the other. This suggests that enough propagules survived between 1998 and 2001 to withstand the differentiating effects of genetic drift.

There are many environmental forces able to exert selective pressures on living organisms. In the case presented here, the different susceptible hosts planted at this location have the potential to select for different genetic characteristics in the *P. capsici* isolates attempting to cause infection. If only a subset of the isolates able to infect cucumbers were able to successfully infect tomatoes then an incomplete sample of the total genetic diversity would be represented by the infecting propagules. Here again we would expect a subset of the genetic diversity recovered in 1998 to be recovered in 2001. In this case, differentiation is not due to random sampling of a small population, but to the non-random nature of the sampling process (eg; *P. capsici* isolates possessing specific genetic characters or constellations of characters are more successful) that occurs under selection. Not only was there no appreciable decrease in the total amount of genetic diversity between 1998 and 2001, but there is no indication that *P. capsici* isolates infecting tomatoes are more similar to each other than they are to isolates recovered from cucumbers. This is illustrated by the genetic similarity tree which showed no increased genetic similarity

(clustering) based on host or year.

We were also interested in the contribution of immigrants to the epidemic in 2001. A previous investigation of the genetic structure of *P. capsici* populations at diverse locations in Michigan suggested that movement among locations was infrequent. Isolates from separate geographical locations were unambiguously more similar to each other, even when comparing fields separated by 1 km (12). If there were significant movement of *P. capsici* propagules into this field, then it is expected that the frequencies of the AFLP markers would differ among years and that novel markers would be introduced in 2001. Marker frequencies did not fluctuate appreciably between years and there were no new AFLP markers detected in 2001. In addition, the frequency of mefenoxam insensitivity remained relatively stable. This suggests that immigration did not contribute significantly to the epidemic occurring in 2001.

In conclusion, it appears that these data support the hypothesis that *P. capsici* survives non-host periods as genetically diverse oospores. Furthermore, this population appears to have maintained its genetic diversity through a 30 month non-host period. How long viable propagules remain in a field following an epidemic is still open to speculation, but it is clear that a typical two year rotation may not ensure against another epidemic. Since it appears that *P. capsici* may remain for extended periods after being introduced and that migration is not a frequent event, then it may be helpful to decipher the mechanisms by which it spreads and develop strategies to limit introduction into new sites.

## **ACKNOWLEDGMENTS**

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Table 1. Estimates of genetic diversity within populations of *Phytophthora capsici* recovered from cucumbers (1998) and tomatoes (2001) planted at the same location in south central Michigan

Year	No. of isolates	Unique isolates <sup>a</sup>	No. of AFLP bands	No. and percent polymorphic bands	Estimated average heterozygosity
1998	57	50	68	42 (62)	0.17
2001	47	43	68	45 (66)	0.17

<sup>a</sup> Total number of isolates with unique multilocus AFLP profiles.

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Table 2: Phenotypic diversity of *Phytophthora capsici* isolates recovered from cucumbers (1998) and tomatoes (2001) planted at the same location in south central Michigan

3 Year	No. of isolates <sup>a</sup>	Compatibility type/mefenoxam sensitivity <sup>b,c</sup>						
			A1/S	A1/IS	A1/I	A2/S	A2/IS	A2/I
4	1998	50	10 (.20)	17 (.34)	2 (.04)	10 (.20)	11 (.22)	-
5	2001	43	7 (.16)	10 (.24)	4 (.09)	12 (.28)	6 (.14)	4 (.09)
6	Total	93	17 (.18)	27 (.30)	6 (.06)	22 (.24)	17 (.18)	4 (.04)

<sup>&</sup>lt;sup>a</sup> Total number of isolates with unique multilocus AFLP profiles.

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b Mefenoxam sensitivity determined by in vitro screening on 100 ppm AI amended media with S = < 30% growth of control (GC), IS = between 30 and 90% GC and I = > 90% GC.

<sup>&</sup>lt;sup>c</sup> Observed numbers are followed by proportion of total sample size in parenthesis.

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Table 3: Results of nested analysis of molecular variance (AMOVA) for *Phytophthora capsici* isolates based on 68 AFLP markers. Variance is partitioned between isolates recovered in 1998 (N = 50) from cucumbers and 2001 (N = 43) from tomatoes at the same location in south central Michigan.

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4	Source of variation <sup>a</sup>	Degrees of freedom	Sum of squares	Variance component	Percentage of variation	P a
5	1998 and 2001					
6	Among populations	1	17.676	0.232	3.40	< 0.0001
7	Within populations	67	618.386	6.578	96.60	

 $<sup>^{</sup>a}$  P = the probability of obtaining a more extreme component estimate by chance alone based on 1000 sampling realizations.

Fig. 1: UPGMA cluster analysis of *Phytophthora capsici* isolates recovered from cucumbers (1998, N = 50) and tomatoes (2001, N = 43) at the same location in south central Michigan based on the Jaccard similarity coefficient using 68 amplified fragment length polymorphism (AFLP) markers. The ratio of isolates recovered in 1998 (cucumber) to the number recovered in 2001 (tomato) within sub-clusters is indicated at major branch points.

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# Worksheet 3-A(10). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress, EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website a websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and thereforesearch has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Endophytes	Study: <u>UNEP 1995, A-73</u>
Section I. Initial Screening on Tech	nical Feasibility of Alternatives
1. Are there any location-specific restrictions that in	hibit the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming of	country
1d Other (Please describe)	

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# Worksheet 3-A(10). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes X No\_\_\_\_\_\_ 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Endophytes 6. Was crop yield measured in the study? Yes No\_\_\_\_\_\_ 7. Describe the effectiveness of the alternative in controlling pests in the study.

8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool?

The use of non-pathogenic endophytes to control Phytophthora capsici is not proven and cannot be considered a				
viable alternative at this time.				
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For EPA	Use Only	
	ID#	

## Worksheet 3-A(11). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

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## **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

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- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

1b. Township caps

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Flooding, Water Management	Study: <u>UNEP 2001, E-74</u>					
Section I. Initial Screening on Technical Feasibility of Alternatives						
1. Are there any location-specific restrictions th	at inhibit the use of this alternative on your site?					
1a. Full use permitted	X					

1c. Alternative not acceptable in consuming country	**************************************
1d. Other (Please describe)	

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## Worksheet 3-A(11). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

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			ID#

## Worksheet 3-A(11)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

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## BACKGROUND

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In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Water Management		The spatiotemporal genetic structure of
		Phytophthora capsici in Michigan and implications
	_	for disease management.

# Section I. Initial Screening on Technical Feasibility of Alternatives

	use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	<del></del>
1d. Other (Please describe)	

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# Worksheet 3-A(11)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

. Is the study on EPA's website	e?	Yes	No	>	<u> </u>		
1a. If not on the EPA w	vebsite, ple	ease attach a copy.					
. Author(s) or researcher(s)	K.H. La	mour					
	M.K. Ha	usbeck	•				
. Publication and Date of Publi	cation	Phytopathology 9	2:681-684, 20	02			
. Location of research study	Michiga	n, USA					
. Name of alternative(s) in stud	y. II more	dian one alternative	,	,,	wish t	o discuss.	
144 4 44	•			-	wish t	o discuss.	
Water Management  Was crop yield measured in the Describe the effectiveness of	he study?	Yesative in controlling p	No_	X udy.			ra capsici
Water Management  . Was crop yield measured in the	he study? the alternatields, cons	Yesative in controlling p	No_ ests in the st	X udy.	d beds,		
Water Management  Was crop yield measured in the secribe the effectiveness of Despite the use of well-drained	he study? the alternatields, cons	Yesative in controlling pervative irrigation, ar	No_ ests in the st	X udy.	d beds,		
. Was crop yield measured in the common of t	he study? the alterna fields, cons	Yes ative in controlling pervative irrigation, ar ply to your situation	No_ ests in the st d planting on r	x udy.	d beds,	Phytophtho	
Water Management  Was crop yield measured in the second of the effectiveness of Despite the use of well-drained is not adequately controlled.  Discuss how the results of the	he study? the alternatields, cons	Yesative in controlling pervative irrigation, ar	No_ ests in the st d planting on r	X udy. aisec	d beds,	Phytophtho	
Water Management  Was crop yield measured in the second of the effectiveness of Despite the use of well-drained is not adequately controlled.  Discuss how the results of the other factors that would affect	he study? the alternatields, cons	Yesative in controlling pervative irrigation, ar	No_ ests in the st d planting on r	X udy. aisec	d beds,	Phytophtho	

## New Frontiers in Plant Disease Losses and Disease Management

# The Spatiotemporal Genetic Structure of *Phytophthora capsici* in Michigan and Implications for Disease Management

K. H. Lamour and M. K. Hausbeck

Department of Plant Pathology, Michigan State University, East Lansing 48824. Accepted for publication 12 February 2002.

Root, crown, and fruit rot caused by *Phytophthora capsici* Leonian is a limiting factor for the production of peppers, tomatoes, and cucurbit crops in Michigan and the United States. Like many species in the genus *Phytophthora*, *P. capsici* has the potential for rapid polycyclic disease development from a limited amount of initial inoculum (6). *P. capsici* produces caducous sporangia that can be spread by wind-blown rain or release 20 to 40 motile zoospores in the presence of free water. The polycyclic phase of disease development is thought to be driven primarily by asexual spore dispersal at a local scale (within and down rows). Sexual reproduction requires both the A1 and A2 compatibility types (CTs) and results in the production of thick-walled oospores. Oospores are thought to serve as the primary survival structure outside of host tissue.

Recommended disease management strategies stress the importance of avoiding excess water in the plant rhizosphere by using well-drained fields, conservative irrigation, and planting on raised beds. Additional recommendations include rotation to nonsusceptible hosts for at least 2 years and the use of fungicides. The phenylamide fungicide (PAF) mefenoxam is a systemic compound with high activity against P. capsici and has been used by growers throughout the United States to control P. capsici. Insensitivity to PAF has been reported for a number of other comycetous organisms (Bremia lactucae, P. infestans, and P. sojae, etc.) and appears to be conferred by a single incompletely dominant gene of major effect (1). Growers in Michigan practicing 2+-year rotation in well-drained fields using an array of fungicidal management tools have experienced significant losses to P. capsici. Michigan is the number one producer of cucumbers for pickling in the United States and it was at the request of grower groups associated with this industry that research into the epidemiology and reproductive biology of P. capsici on cucurbit hosts was initiated.

Although many researchers cite oospores as the most likely propagule for survival outside of host tissue, there have been very few investigations specifically aimed at determining the impact of sexual reproduction in natural populations of *P. capsici*. Our hypothesis was that the sexual stage may play an important role not only in survival but also in the adaptation of *P. capsici* populations to environmental stresses (e.g., fungicides). Our goal was to perform a comprehensive investigation of the phenotypic and genetic diversity present in *P. capsici* populations from the major vegetable production regions of Michigan, with the implicit intention of addressing questions concerning epidemiology, repro-

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ductive biology, and the durability of currently recommended management strategies.

#### METHODOLOGY

Isolate collection and maintenance. Sampling of diseased fields began at the end of the 1997 growing season and continued through September 2000. In all cases, fields were sampled on a grid with quadrants varying from 40 m² to 12 km².—A limited number of isolates were collected in 1997. In 1998, the strategy was to collect as many samples from as many fields as possible. This strategy was modified in 1999 and 2000 to focus on specific fields. Isolations from diseased plants were made onto selective media and single zoospore cultures were generated according to standard single sporing techniques (3). Isolates were placed into long-term storage (15°C) using a hemp seed/sterile water technique.

Phenotypic characterization. Single zoospore isolates were screened for CT using known A1 and A2 isolates. In vitro screening techniques published for other Phytophthora species for assessing sensitivity to mefenoxam were compared and a novel. simple, high dose screen using 100 ppm of mefenoxam-amended V8 agar was found to separate field isolates into three modal distributions that appeared consistent with the expectations of a single incompletely dominant gene governing mefenoxam insensitivity (e.g., sensitive, intermediately sensitive, and fully insensitive). These putative mefenoxam sensitivity (MS) groupings were tested by performing a series of crosses and testing whether the observed progeny sets met the expectations for Mendelian inheritance of a single incompletely dominant gene controlling insensitivity to mefenoxam. Sexual crosses were conducted on unclarified V8 agar plates and incubated for 3 months in the dark. Individual germinated oospores were recovered after 3 months using previously published techniques (2).

The efficacy of this in vitro mefenoxam screening technique was further tested in pumpkin seedlings using progeny from a cross between parents intermediately sensitive to mefenoxam. Nine isolates from each of the three MS categories were screened for pathogenicity on untreated seedlings. Single sensitive, intermediately sensitive, and fully insensitive isolates were then placed onto the unwounded surface of plants treated with either a field rate of mefenoxam, three times the field rate, or distilled water. Lesion diameters on seedling stems were measured after 4 days.

Genetic characterization. Single zoospore isolates were grown in antibiotic-amended V8 broth for 3 days at room temperature. Mycelial mats were washed, frozen, lyophilized, and ground with a sterile mortar and pestle. DNA was extracted with either a Qiagen Dneasy extraction kit (Qiagen, Valencia, CA) or via a cetyltrimethylammonium bromide (CTAB) procedure. A variety

of methods for generating molecular markers were tested for efficacy including isozyme, random amplified polymorphic DNA, and amplified fragment length polymorphism (AFLP). The AFLP technique resulted in a large number of reproducible markers and was chosen to characterize samples of P. capsici from Michigan. The AFLP technique involves cutting genomic DNA with moderately rare cutting (EcoRI) and frequent cutting (MseI) restriction enzymes, while concomitantly ligating synthetic adaptor fragments of DNA to the sticky ends created by the restriction enzymes (7). The result is a large number of DNA fragments that have ends with known DNA sequences. Amplification of fragment subsets (termed fingerprints) can be accomplished using polymerase chain reaction (PCR) primers complementary to the adaptor sequences with additional "selective" nucleotides. Changing the amount and type of selective nucleotides results in different subsets or fingerprints. Stringent PCR cycling parameters (touchdown technique) are used to ensure the fidelity of the reaction. For the analysis summarized here, adaptor sequences and fluorescent labeled selective primers were purchased as a kit through Perkin-Elmer ABI (Applied Biosystems, Foster City, CA). Using this system, AFLP fragments were resolved on a polyacrylamide gel by an ABI 377 gene sequencer. Fluorescent labels were excited by a laser and band emissions were analyzed in the form of an electropherogram where peaks represent individual bands. The sizing of fragments was particularly robust because a DNA ladder was loaded with every sample into the gel. To test for the reproducibility of fingerprints, DNA was extracted from a single isolate on three separate occasions approximately 3 months apart and subjected to the aforementioned protocol.

Data analysis. Isolates with identical multilocus AFLP fingerprints were considered to be members of the same clonal lineage and only a single representative was used for analysis. Because AFLP markers can only be scored confidently for presence (1) or absence (0), allele frequencies were estimated based on the assumption that populations under investigation meet the criterion for Hardy-Weinberg equilibrium, and that loci have only one "present" allele. The term population refers to all samples taken from a single field during a single year.

Genetic diversity within single populations was assessed by calculating the average number of polymorphic bands and estimating the average heterozygosity. Fixation indices were calculated according to methods of Weir and Cockerham (8) for populations from the same site over multiple years and among populations in Michigan using the program tools for population genetic analysis (TFPGA) (M. P. Miller, Northern Arizona University, Flagstaff). Confidence intervals for F statistics at the 95% confidence level were generated by bootstrapping at 1,000 iterations. The program NTSYS-pc version 2.02k (Exeter Software, Setauket, NY) was used to construct a similarity matrix from the presence/absence (1/0) data. Cluster analysis using the unweighted pair group with arithmetic averages (UPGMA) method was performed on the matrix and a tree was generated to give a visual representation of isolate similarity. Excoffier's ARLEQUIN program (L. Excoffier, University of Geneva) was used to assess population differentiation using a phenetic approach termed analysis of molecular variance (AMOVA), which allows for total genetic variation to be partitioned within and among populations using a classical analysis of variance (ANOVA).

## **RESULTS**

Phenotypic results. Five isolates were recovered in 1997 from five different farms (four A1 and one A2 CT). One isolate was fully insensitive to mefenoxam, whereas the other four were fully sensitive. These findings prompted the extensive sampling conducted in 1998 in which 523 isolates (473 from cucurbits and 30 from bell pepper) were collected from 14 farms. A frequency histogram plotting percent growth of control on 100 ppm of

mefenoxam-amended media versus number of isolates revealed a trimodal distribution (3). Putative MS categories were assigned based on these groupings with sensitive (S) <30% growth of control, intermediately sensitive (IS) between 30 and 90% growth of control, and insensitive (I) >90% growth of control. In vitro crosses between isolates representative of the different putative sensitivity categories (S  $\times$  S, I  $\times$  S, IS  $\times$  S, and IS  $\times$  IS) resulted in progeny sets not significantly different than expected for insensitivity inherited as a single incompletely dominant gene unlinked to CT (P = 0.05) (3). In 1998, 55% of the isolates were sensitive to mefenoxam, 32% were intermediately sensitive, and 13% were fully insensitive to mefenoxam. A1 and A2 CTs were recovered in a ratio of approximately 1:1 in 8 of the 14 farms. Oospores were detected in naturally diseased cucurbit fruit from four farms, and 223 oospore progeny were recovered and germinated from a single diseased cucumber. All six possible MS × CT combinations were detected in this naturally occurring oospore progeny set (3).

In planta studies using sensitive, intermediately sensitive, and fully insensitive *P. capsici* isolates supported the in vitro screening categories, with sensitive isolates causing no disease on mefenoxam-treated plants, intermediately sensitive isolates being slowed by mefenoxam, and fully insensitive isolates showing no difference in the ability to colonize host tissue between treated and untreated plants at three times the field rate. All the progeny isolates were pathogenic on untreated pumpkin plants (K. H. Lamour and M. K. Hausbeck, *unpublished data*).

Sixty-three mefenoxam insensitive (18% intermediate and 82% fully insensitive) isolates were recovered from a single southwest Michigan field in 1998. Field experiments were conducted in this field during 1999 and 2000, testing alternative cultural control strategies, and no mefenoxam was applied. Two hundred isolates were recovered from this site over the course of the 1999 season and 34 isolates at the beginning of the 2000 season. Of the 200 isolates recovered in 1999 from this field, 141 had unique AFLP genotypes. Seventy percent of these were fully insensitive to mefenoxam, 28% were intermediately sensitive, and 2% were sensitive. In 2000, 15% of the isolates were intermediately sensitive and 85% were fully insensitive. A single fully insensitive clonal lineage rose in frequency over the course of the 1999 season and comprised 20% of the total number of samples recovered (4).

During 1999 and 2000, approximately 2,500 isolates were recovered from farms in Michigan. Both the A1 and A2 CTs were present in every field sampled, and mefenoxam insensitivity was detected in the majority of farms that had a history of mefenoxam use.

Genetic results. Nine populations from the four major vegetable production areas of Michigan were analyzed with the AFLP procedure (N = 641). AFLP analysis resolved a total of 94 clearly discernable markers when considering all the isolates together. No single isolate or group of isolates from a single location contained all 94 markers. The total number of AFLP loci in a single population ranged from 68 to 80. Seventeen (18%) fragments were fixed for the present state across all populations, 12 (13%) fragments were polymorphic in all populations, and 65 (69%) were fixed for presence or absence in some populations and polymorphic in others. The number of polymorphic bands within a single population ranged from 37 to 46 with estimated heterozygosities ranging from 0.18 to 0.22. Clonal reproduction was significant within single fields over the course of the growing season. For example, genotypic diversity in a single field ranged from 100% at the beginning of the growing season (seedling stage) to <30% at the time cucurbit fruit were ready for harvest (4). When considering all nine populations, genotypic diversity ranged from 42 to 96% with an average of 74% of the isolates in any sample set having unique genotypes. Although clonal reproduction was significant within single fields within years, no clones were recovered from single fields between years or among fields separated by at least 1 km. Fixation indices ( $\phi_{ST}$ ) between the

populations sampled on consecutive years were very close to zero, indicating that gene diversity was not measurably impacted by genetic drift (5). The overall estimated  $\phi_{ST}$  for populations from different locations was 0.35, indicating that approximately 35% of the total genetic diversity present in Michigan *P. capsici* populations is found among populations and 65% is found within any one population. AMOVA partitioned genetic diversity among (40%) and within (60%) populations. The similarity tree based on UPGMA cluster analysis clearly showed that isolates from the

same site sampled over years branched from the same node, with no clustering of isolates based on the year of sampling. Cluster analysis also clearly showed that populations separated geographically branched from population-specific nodes (5).

## **DISCUSSION**

During the past 10 years, Michigan has experienced a steady increase in the incidence of root, fruit, and crown rot on cucurbits

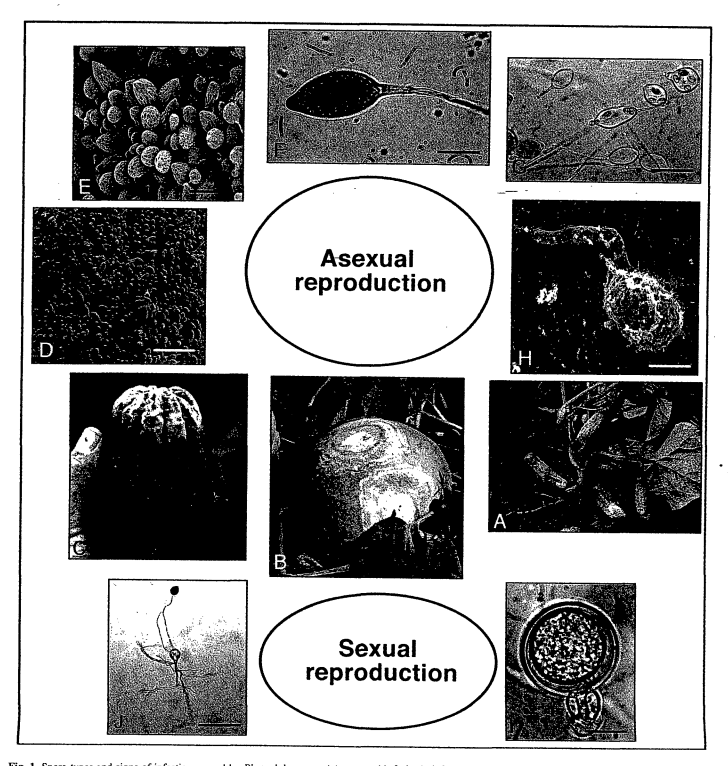


Fig. 1. Spore types and signs of infection caused by *Phytophthora capsici* on cucurbit fruit: A, infected cucumber, B, pumpkin, and C, acorn squash fruit. D, Scanning electron microscope (SEM) photo of an infected cucumber showing tufts of sporangia produced on the surface of the fruit (Bar = 30  $\mu$ m). E, Close-up of a single tuft of sporangia (Bar = 30  $\mu$ m). F, Typical papillate sporangium with a long pedicel (Bar = 20  $\mu$ m). G, Zoospores exiting sporangia after immersion in water (Bar = 50  $\mu$ m). H, SEM photo of a single encysted zoospore that germinated and directly penetrated the epidermis of a cucumber fruit (Bar = 4  $\mu$ m). I, Typical amphigynous oospore (Bar = 10  $\mu$ m). J, A germinating oospore with multiple germ tubes and a terminal sporangium (Bar = 100  $\mu$ m).

caused by *P. capsici*. Rotation to nonsusceptible hosts, in conjunction with cultural and chemical control strategies, have not provided economic control. Correspondence with other vegetable pathologists suggests that this phenomenon is not confined to Michigan, and a similar increase in control failures due to blight by *P. capsici* is being reported throughout the United States.

Investigation of the inheritance of MS demonstrated that MS is inherited as a single incompletely dominant gene unlinked to CT. In 1998, all six possible MS × CT combinations were present in single fields and insensitivity to mefenoxam was common in Michigan. Typical amphigynous oospores were observed in *P. capsici*-infected cucurbit fruit from multiple locations, and oospore progeny from a single naturally infected fruit showed segregation for MS and CT. These findings strongly support the hypothesis that sexual reproduction is occurring in the field, and also suggest that sexual recombination may directly generate progeny fully insensitive to mefenoxam. Tracking a single mefenoxam insensitive population over 2 years in the absence of mefenoxam selection pressure suggests that costs associated with mefenoxam insensitivity are minimal.

Estimates of average heterozygosity and polymorphism indicate surprisingly high levels of gene and genotypic diversity in all the populations of P. capsici analyzed. Tracking a single population through an entire growing season showed that asexual reproduction plays a significant role in disease development within a single season. Sampling single fields over consecutive years suggested that clones do not survive Michigan winters and that oospores are the primary survival propagule. Estimation of fixation indices for samples from the same site over consecutive years suggested that there was not a significant reduction in genetic diversity between growing seasons. This implies that populations are large enough to withstand dramatic effects of genetic drift. Cluster analysis revealed unambiguous groups corresponding to geographical locations with regional populations showing more similarity overall than populations from different regions. Population pairwise fixation indices corroborated this finding. The estimated overall fixation index and AMOVA are in agreement with both, suggesting that most (approx 60%) of the total genetic variability in Michigan is found within any one population, but that a relatively large component (40%) of genetic variability is found among populations.

Recommendations based on our findings are as follows: (i) the fungicide mefenoxam may be of limited usefulness because insensitivity appears to be selected for rapidly and is unlikely to decrease when mefenoxam selection pressure is removed; (ii) fields with epidemics are likely to harbor oospores for an extended amount of time (at least 5 years), and this factor must be considered before replanting to susceptible hosts; and (iii) factors that may contribute to the introduction of *P. capsici* into uninfested fields (e.g., drainage ditches between farms, irrigation ponds, and the dumping of culls) need to be considered and if possible avoided, because once an epidemic is established we have found no evidence that the population will become extinct in an agriculturally meaningful time period.

From an evolutionary perspective, it is clear that P. capsici has successfully colonized a number of geographical locations in

Michigan and that each of the populations sampled thus far have similarly high levels of genetic variability. The genetic stability of single populations over multiple years, the high fixation indices between even geographically close populations (1 km), and the clear structuring based on UPGMA cluster analysis all suggest that long-distance dispersal of inoculum is not common and that geographically isolated populations are also genetically isolated. It appears that the sexual stage of the P. capsici life cycle plays a significant role in survival as well as maintaining both genic and genotypic diversity, and has likely played a key role in the evolution of mefenoxam insensitivity. The combination of high levels of genetic variability, thick-walled oospores, and polycyclic asexual disease development make P. capsici a formidable pathogen (Fig. 1). This work underscores the need for management strategies aimed at preventing the spread of P. capsici to uninfested field sites and suggests that management strategies aimed at limiting spread within a single season may be the only option for growers with P. capsici-infested fields.

#### ACKNOWLEDGMENTS

This work was funded by the Michigan Agricultural Experiment Station, Michigan State University Extension, Michigan Department of Agriculture, Michigan Farm Bureau (GREEN cooperative), Pickle and Pepper Research Committee, Pickle Packers International, Inc., and the Pickle Seed Research Fund, Pickle Packers International. We thank M. Bour, C. Hunter, J. Jabara, P. Tumbalam, E. Webster, and J. Woodworth for competent laboratory assistance. K. Lamour thanks his Ph.D. committee members A. Jarosz, R. Hammerschmidt, and F. Trail for guidance and extends sincere thanks to M. Hausbeck for fulfilling her role as mentor in an exemplary manner.

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			ID#	

# Worksheet 3-A(12). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by\_crep-and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: General IPM	Study: UNEP 1998, B-91, B-94, B-288

# Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the u	use of this alternative on your s	ite?
--	-----------------------------------	------

1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	

For EPA Use Only	
ID#	

# Worksheet 3-A(12). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

1. Is the study on EPA's website?	Yes <u>X</u> No
1a. If not on the EPA website, pl	lease attach a copy.
2. Author(s) or researcher(s)	
3. Publication and Date of Publication	
4. Location of research study	
5. Name of alternative(s) in study. If more  General IPM	than one alternative, list the ones you wish to discuss.
6. Was crop yield measured in the study?	? Yes No
7. Describe the effectiveness of the altern	native in controlling pests in the study.
<u> </u>	
3. Discuss how the results of the study ap other factors that would affect your ado	pply to your situation. Would you expect similar results? Are there option of this tool?
I would not expect similar results because	Michigan cucurbit growers use extensive IPM practices, but have severe
	million to Compare the IDM and Compare that
disease due to Phytophthora crown and from	ruit rot. Some of the IPM practices that growers use include crop rotation,
	ruit rot. Some of the IPM practices that growers use include crop rotation, rungicide sprays. These practices, even when used in combination, have

# Worksheet 3-A(12)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: General IPM .	Study: Alternatives to methyl bromide for vegetable		
	covered crops in Morocco		
Section I. Initial Screening on	Technical Feasibility of Alternatives		
1. Are there any location-specific restriction	ns that inhibit the use of this alternative on your site?		
1a. Full use permitted	X		
1b. Township caps			
1c. Alternative not acceptable in con-	suming country		
1d. Other (Please describe)			

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# Worksheet 3-A(12)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

1. Is the study on EPA's website?	Yes X No
1a. If not on the EPA website, p	lease attach a copy.
2. Author(s) or researcher(s)	
Publication and Date of Publication	
Location of research study	
5. Name of alternative(s) in study. If more General IPM	e than one alternative, list the ones you wish to discuss.
5. Was crop yield measured in the study	? Yes No
	native in controlling pests in the study.
7. Describe the effectiveness of the altern	
7. Describe the effectiveness of the altern	
	pply to your situation. Would you expect similar results? Are there option of this tool?
3. Discuss how the results of the study a other factors that would affect your ad This study does not apply to the Michigan	pply to your situation. Would you expect similar results? Are there

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## Worksheet 3-A(12)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

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- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

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Use additional pages as needed.

Alternative: General IPM		The spatiotemporal genetic structure of
	-	Phytophthora capsici in Michigan and implications
		for disease management.

## Section I. Initial Screening on Technical Feasibility of Alternatives

1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	

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	ID#	

# Worksheet 3-A(12)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

Is the study on EPA's website	?	Yes	No	<u>X</u>	- Ji
1a. If not on the EPA w	ebsite, plea	se attach a copy.			-
Author(s) or researcher(s)	K.H. Lamo	our			
	M.K. Haus	beck			
Publication and Date of Public	cation	Phytopathology 9	2:681-684, 2002	2	
Location of research study	Michigan,	USA			
Was crop yield measured in the Describe the effectiveness of	the alternati	Yesve in controlling p	No	X	ppagement planting
Was crop yield measured in the Describe the effectiveness of Growers in Michigan follow reco	he study? the alternationmended di	Yesve in controlling p	No  ests in the stud t strategies, incl	X dy. uding water ma	anagement, planting
Was crop yield measured in the Describe the effectiveness of Growers in Michigan follow recommends on raised beds, rotation to nons	he study? the alternationmended disusceptible ho	Yesve in controlling psease management osts and the use of	No	X  dy.  uding water mastill suffer losse	anagement, planting
Was crop yield measured in the Describe the effectiveness of Growers in Michigan follow reco	he study? the alternationmended disusceptible ho	Yesve in controlling p	No_ nests in the stud t strategies, incl fungicides, and	X  dy.  uding water ma	anagement, planting

## New Frontiers in Plant Disease Losses and Disease Management

# The Spatiotemporal Genetic Structure of *Phytophthora capsici* in Michigan and Implications for Disease Management

K. H. Lamour and M. K. Hausbeck

Department of Plant Pathology, Michigan State University, East Lansing 48824. Accepted for publication 12 February 2002.

Root, crown, and fruit rot caused by *Phytophthora capsici* Leonian is a limiting factor for the production of peppers, tomatoes, and cucurbit crops in Michigan and the United States. Like many species in the genus *Phytophthora*, *P. capsici* has the potential for rapid polycyclic disease development from a limited amount of initial inoculum (6). *P. capsici* produces caducous sporangia that can be spread by wind-blown rain or release 20 to 40 motile zoospores in the presence of free water. The polycyclic phase of disease development is thought to be driven primarily by asexual spore dispersal at a local scale (within and down rows). Sexual reproduction requires both the A1 and A2 compatibility types (CTs) and results in the production of thick-walled oospores. Oospores are thought to serve as the primary survival structure outside of host tissue.

Recommended disease management strategies stress the importance of avoiding excess water in the plant rhizosphere by using well-drained fields, conservative irrigation, and planting on raised beds. Additional recommendations include rotation to nonsusceptible hosts for at least 2 years and the use of fungicides. The phenylamide fungicide (PAF) mefenoxam is a systemic compound with high activity against P. capsici and has been used by growers throughout the United States to control P. capsici. Insensitivity to PAF has been reported for a number of other oomycetous organisms (Bremia lactucae, P. infestans, and P. sojae, etc.) and appears to be conferred by a single incompletely dominant gene of major effect (1). Growers in Michigan practicing 2+-year rotation in well-drained fields using an array of fungicidal management tools have experienced significant losses to P. capsici. Michigan is the number one producer of cucumbers for pickling in the United States and it was at the request of grower groups associated with this industry that research into the epidemiology and reproductive biology of P. capsici on cucurbit hosts was initiated.

Although many researchers cite oospores as the most likely propagule for survival outside of host tissue, there have been very few investigations specifically aimed at determining the impact of sexual reproduction in natural populations of *P. capsici*. Our hypothesis was that the sexual stage may play an important role not only in survival but also in the adaptation of *P. capsici* populations to environmental stresses (e.g., fungicides). Our goal was to perform a comprehensive investigation of the phenotypic and genetic diversity present in *P. capsici* populations from the major vegetable production regions of Michigan, with the implicit intention of addressing questions concerning epidemiology, repro-

Corresponding author: K. H. Lamour; E-mail address: lamourku@msu.edu

ductive biology, and the durability of currently recommended management strategies.

## METHODOLOGY

Isolate collection and maintenance. Sampling of diseased fields began at the end of the 1997 growing season and continued through September 2000. In all cases, fields were sampled on a grid with quadrants varying from 40 m² to 12 km². A limited number of isolates were collected in 1997. In 1998, the strategy was to collect as many samples from as many fields as possible. This strategy was modified in 1999 and 2000 to focus on specific fields. Isolations from diseased plants were made onto selective media and single zoospore cultures were generated according to standard single sporing techniques (3). Isolates were placed into long-term storage (15°C) using a hemp seed/sterile water technique.

Phenotypic characterization. Single zoospore isolates were screened for CT using known A1 and A2 isolates. In vitro screening techniques published for other Phytophthora species for assessing sensitivity to mefenoxam were compared and a novel, simple, high dose screen using 100 ppm of mefenoxam-amended V8 agar was found to separate field isolates into three modal distributions that appeared consistent with the expectations of a single incompletely dominant gene governing mefenoxam insensitivity (e.g., sensitive, intermediately sensitive, and fully insensitive). These putative mefenoxam sensitivity (MS) groupings were tested by performing a series of crosses and testing whether the observed progeny sets met the expectations for Mendelian inheritance of a single incompletely dominant gene controlling insensitivity to mefenoxam. Sexual crosses were conducted on unclarified V8 agar plates and incubated for 3 months in the dark. Individual germinated oospores were recovered after 3 months using previously published techniques (2).

The efficacy of this in vitro mefenoxam screening technique was further tested in pumpkin seedlings using progeny from a cross between parents intermediately sensitive to mefenoxam. Nine isolates from each of the three MS categories were screened for pathogenicity on untreated seedlings. Single sensitive, intermediately sensitive, and fully insensitive isolates were then placed onto the unwounded surface of plants treated with either a field rate of mefenoxam, three times the field rate, or distilled water. Lesion diameters on seedling stems were measured after 4 days.

Genetic characterization. Single zoospore isolates were grown in antibiotic-amended V8 broth for 3 days at room temperature. Mycelial mats were washed, frozen, lyophilized, and ground with a sterile mortar and pestle. DNA was extracted with either a Qiagen Dneasy extraction kit (Qiagen, Valencia, CA) or via a cetyltrimethylammonium bromide (CTAB) procedure. A variety

of methods for generating molecular markers were tested for efficacy including isozyme, random amplified polymorphic DNA, and amplified fragment length polymorphism (AFLP). The AFLP technique resulted in a large number of reproducible markers and was chosen to characterize samples of P. capsici from Michigan. The AFLP technique involves cutting genomic DNA with moderately rare cutting (EcoRI) and frequent cutting (MseI) restriction enzymes, while concomitantly ligating synthetic adaptor fragments of DNA to the sticky ends created by the restriction enzymes (7). The result is a large number of DNA fragments that have ends with known DNA sequences. Amplification of fragment subsets (termed fingerprints) can be accomplished using polymerase chain reaction (PCR) primers complementary to the adaptor sequences with additional "selective" nucleotides. Changing the amount and type of selective nucleotides results in different subsets or fingerprints. Stringent PCR cycling parameters (touchdown technique) are used to ensure the fidelity of the reaction. For the analysis summarized here, adaptor sequences and fluorescent labeled selective primers were purchased as a kit through Perkin-Elmer ABI (Applied Biosystems, Foster City, CA). Using this system, AFLP fragments were resolved on a polyacrylamide gel by an ABI 377 gene sequencer. Fluorescent labels were excited by a laser and band emissions were analyzed in the form of an electropherogram where peaks represent individual bands. The sizing of fragments was particularly robust because a DNA ladder was loaded with every sample into the gel. To test for the reproducibility of fingerprints, DNA was extracted from a single isolate on three separate occasions approximately 3 months apart and subjected to the aforementioned protocol.

Data analysis. Isolates with identical multilocus AFLP fingerprints were considered to be members of the same clonal lineage and only a single representative was used for analysis. Because AFLP markers can only be scored confidently for presence (1) or absence (0), allele frequencies were estimated based on the assumption that populations under investigation meet the criterion for Hardy-Weinberg equilibrium, and that loci have only one "present" allele. The term population refers to all samples taken from a single field during a single year.

Genetic diversity within single populations was assessed by calculating the average number of polymorphic bands and estimating the average heterozygosity. Fixation indices were calculated according to methods of Weir and Cockerham (8) for populations from the same site over multiple years and among populations in Michigan using the program tools for population genetic analysis (TFPGA) (M. P. Miller, Northern Arizona University, Flagstaff). Confidence intervals for F statistics at the 95% confidence level were generated by bootstrapping at 1,000 iterations. The program NTSYS-pc version 2.02k (Exeter Software, Setauket, NY) was used to construct a similarity matrix from the presence/absence (1/0) data. Cluster analysis using the unweighted pair group with arithmetic averages (UPGMA) method was performed on the matrix and a tree was generated to give a visual representation of isolate similarity. Excoffier's ARLEQUIN program (L. Excoffier, University of Geneva) was used to assess population differentiation using a phenetic approach termed analysis of molecular variance (AMOVA), which allows for total genetic variation to be partitioned within and among populations using a classical analysis of variance (ANOVA).

## **RESULTS**

Phenotypic results. Five isolates were recovered in 1997 from five different farms (four A1 and one A2 CT). One isolate was fully insensitive to mefenoxam, whereas the other four were fully sensitive. These findings prompted the extensive sampling conducted in 1998 in which 523 isolates (473 from cucurbits and 30 from bell pepper) were collected from 14 farms. A frequency histogram plotting percent growth of control on 100 ppm of

mefenoxam-amended media versus number of isolates revealed a trimodal distribution (3). Putative MS categories were assigned based on these groupings with sensitive (S) <30% growth of control, intermediately sensitive (IS) between 30 and 90% growth of control, and insensitive (I) >90% growth of control. In vitro crosses between isolates representative of the different putative sensitivity categories (S  $\times$  S, I  $\times$  S, IS  $\times$  S, and IS  $\times$  IS) resulted in progeny sets not significantly different than expected for insensitivity inherited as a single incompletely dominant gene unlinked to CT (P = 0.05) (3). In 1998, 55% of the isolates were sensitive to mefenoxam, 32% were intermediately sensitive, and 13% were fully insensitive to mefenoxam. A1 and A2 CTs were recovered in a ratio of approximately 1:1 in 8 of the 14 farms. Oospores were detected in naturally diseased cucurbit fruit from four farms, and 223 oospore progeny were recovered and germinated from a single diseased cucumber. All six possible MS × CT combinations were detected in this naturally occurring oospore progeny set (3).

In planta studies using sensitive, intermediately sensitive, and fully insensitive *P. capsici* isolates supported the in vitro screening categories, with sensitive isolates causing no disease on mefenoxam-treated plants, intermediately sensitive isolates being slowed by mefenoxam, and fully insensitive isolates showing no difference in the ability to colonize host tissue between treated and untreated plants at three times the field rate. All the progeny isolates were pathogenic on untreated pumpkin plants (K. H. Lamour and M. K. Hausbeck, *unpublished data*).

Sixty-three mefenoxam insensitive (18% intermediate and 82% fully insensitive) isolates were recovered from a single southwest Michigan field in 1998. Field experiments were conducted in this field during 1999 and 2000, testing alternative cultural control strategies, and no mefenoxam was applied. Two hundred isolates were recovered from this site over the course of the 1999 season and 34 isolates at the beginning of the 2000 season. Of the 200 isolates recovered in 1999 from this field, 141 had unique AFLP genotypes. Seventy percent of these were fully insensitive to mefenoxam, 28% were intermediately sensitive, and 2% were sensitive. In 2000, 15% of the isolates were intermediately sensitive and 85% were fully insensitive. A single fully insensitive clonal lineage rose in frequency over the course of the 1999 season and comprised 20% of the total number of samples recovered (4).

During 1999 and 2000, approximately 2,500 isolates were recovered from farms in Michigan. Both the A1 and A2 CTs were present in every field sampled, and mefenoxam insensitivity was detected in the majority of farms that had a history of mefenoxam use.

Genetic results. Nine populations from the four major vegetable production areas of Michigan were analyzed with the AFLP procedure (N = 641). AFLP analysis resolved a total of 94 clearly discernable markers when considering all the isolates together. No single isolate or group of isolates from a single location contained all 94 markers. The total number of AFLP loci in a single population ranged from 68 to 80. Seventeen (18%) fragments were fixed for the present state across all populations, 12 (13%) fragments were polymorphic in all populations, and 65 (69%) were fixed for presence or absence in some populations and polymorphic in others. The number of polymorphic bands within a single population ranged from 37 to 46 with estimated heterozygosities ranging from 0.18 to 0.22. Clonal reproduction was significant within single fields over the course of the growing season. For example, genotypic diversity in a single field ranged from 100% at the beginning of the growing season (seedling stage) to <30% at the time cucurbit fruit were ready for harvest (4). When considering all nine populations, genotypic diversity ranged from 42 to 96% with an average of 74% of the isolates in any sample set having unique genotypes. Although clonal reproduction was significant within single fields within years, no clones were recovered from single fields between years or among fields separated by at least 1 km. Fixation indices ( $\phi_{ST}$ ) between the

populations sampled on consecutive years were very close to zero, indicating that gene diversity was not measurably impacted by genetic drift (5). The overall estimated  $\phi_{ST}$  for populations from different locations was 0.35, indicating that approximately 35% of the total genetic diversity present in Michigan *P. capsici* populations is found among populations and 65% is found within any one population. AMOVA partitioned genetic diversity among (40%) and within (60%) populations. The similarity tree based on UPGMA cluster analysis clearly showed that isolates from the

same site sampled over years branched from the same node, with no clustering of isolates based on the year of sampling. Cluster analysis also clearly showed that populations separated geographically branched from population-specific nodes (5).

## DISCUSSION

During the past 10 years, Michigan has experienced a steady increase in the incidence of root, fruit, and crown rot on cucurbits

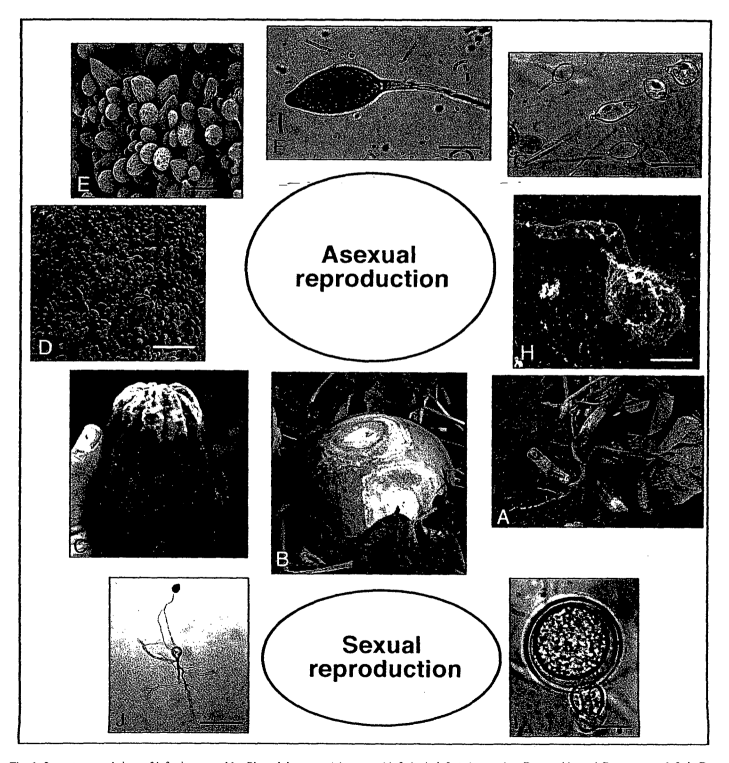


Fig. 1. Spore types and signs of infection caused by *Phytophthora capsici* on cucurbit fruit: A, infected cucumber, B, pumpkin, and C, acorn squash fruit. D, Scanning electron microscope (SEM) photo of an infected cucumber showing tufts of sporangia produced on the surface of the fruit (Bar = 300  $\mu$ m). E, Close-up of a single tuft of sporangia (Bar = 30  $\mu$ m). F, Typical papillate sporangium with a long pedicel (Bar = 20  $\mu$ m). G, Zoospores exiting sporangia after immersion in water (Bar = 50  $\mu$ m). H, SEM photo of a single encysted zoospore that germinated and directly penetrated the epidermis of a cucumber fruit (Bar = 4  $\mu$ m). I, Typical amphigynous oospore (Bar = 10  $\mu$ m). J, A germinating oospore with multiple germ tubes and a terminal sporangium (Bar = 100  $\mu$ m).

caused by *P. capsici*. Rotation to nonsusceptible hosts, in conjunction with cultural and chemical control strategies, have not provided economic control. Correspondence with other vegetable pathologists suggests that this phenomenon is not confined to Michigan, and a similar increase in control failures due to blight by *P. capsici* is being reported throughout the United States.

Investigation of the inheritance of MS demonstrated that MS is inherited as a single incompletely dominant gene unlinked to CT. In 1998, all six possible MS × CT combinations were present in single fields and insensitivity to mefenoxam was common in Michigan. Typical amphigynous oospores were observed in P. capsici-infected cucurbit fruit from multiple locations, and oospore progeny from a single naturally infected fruit showed segregation for MS and CT. These findings strongly support the hypothesis that sexual reproduction is occurring in the field, and also suggest that sexual recombination may directly generate progeny fully insensitive to mefenoxam. Tracking a single mefenoxam insensitive population over 2 years in the absence of mefenoxam selection pressure suggests that costs associated with mefenoxam insensitivity are minimal.

Estimates of average heterozygosity and polymorphism indicate surprisingly high levels of gene and genotypic diversity in all the populations of P. capsici analyzed. Tracking a single population through an entire growing season showed that asexual reproduction plays a significant role in disease development within a single season. Sampling single fields over consecutive years suggested that clones do not survive Michigan winters and that oospores are the primary survival propagule. Estimation of fixation indices for samples from the same site over consecutive years suggested that there was not a significant reduction in genetic diversity between growing seasons. This implies that populations are large enough to withstand dramatic effects of genetic drift. Cluster analysis revealed unambiguous groups corresponding to geographical locations with regional populations showing more similarity overall than populations from different regions. Population pairwise fixation indices corroborated this finding. The estimated overall fixation index and AMOVA are in agreement with both, suggesting that most (approx 60%) of the total genetic variability in Michigan is found within any one population, but that a relatively large component (40%) of genetic variability is found among populations.

Recommendations based on our findings are as follows: (i) the fungicide mefenoxam may be of limited usefulness because insensitivity appears to be selected for rapidly and is unlikely to decrease when mefenoxam selection pressure is removed; (ii) fields with epidemics are likely to harbor oospores for an extended amount of time (at least 5 years), and this factor must be considered before replanting to susceptible hosts; and (iii) factors that may contribute to the introduction of *P. capsici* into uninfested fields (e.g., drainage ditches between farms, irrigation ponds, and the dumping of culls) need to be considered and if possible avoided, because once an epidemic is established we have found no evidence that the population will become extinct in an agriculturally meaningful time period.

From an evolutionary perspective, it is clear that P. capsici has successfully colonized a number of geographical locations in

Michigan and that each of the populations sampled thus far have similarly high levels of genetic variability. The genetic stability of single populations over multiple years, the high fixation indices between even geographically close populations (1 km), and the clear structuring based on UPGMA cluster analysis all suggest that long-distance dispersal of inoculum is not common and that geographically isolated populations are also genetically isolated. It appears that the sexual stage of the P. capsici life cycle plays a significant role in survival as well as maintaining both genic and genotypic diversity, and has likely played a key role in the evolution of mefenoxam insensitivity. The combination of high levels of genetic variability, thick-walled oospores, and polycyclic asexual disease development make P. capsici a formidable pathogen (Fig. 1). This work underscores the need for management strategies aimed at preventing the spread of P. capsici to uninfested field sites and suggests that management strategies aimed at limiting spread within a single season may be the only option for growers with P. capsici-infested fields.

## **ACKNOWLEDGMENTS**

This work was funded by the Michigan Agricultural Experiment Station, Michigan State University Extension, Michigan Department of Agriculture, Michigan Farm Bureau (GREEEN cooperative), Pickle and Pepper Research Committee, Pickle Packers International, Inc., and the Pickle Seed Research Fund, Pickle Packers International. We thank M. Bour, C. Hunter, J. Jabara, P. Tumbalam, E. Webster, and J. Woodworth for competent laboratory assistance. K. Lamour thanks his Ph.D. committee members A. Jarosz, R. Hammerschmidt, and F. Trail for guidance and extends sincere thanks to M. Hausbeck for fulfilling her role as mentor in an exemplary manner.

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## Worksheet 3-A(12)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need. For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b). When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8. Summarize each of the research studies you cite in the Research Summary Worksheet. If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed. BACKGROUND EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996. There are three major ways you can provide the Agency with proof of your investigative work. (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates. application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website. The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area. In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area. Use additional pages as needed. Alternative: General IPM . Study: Alternatives for methyl bromide on cucurbits, 2002. Section I. Initial Screening on Technical Feasibility of Alternatives 1. Are there any location-specific restrictions that inhibit the use of this alternative on your site? 1a. Full use permitted 1b. Township caps 1c. Alternative not acceptable in consuming country 1d. Other (Please describe)

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# Worksheet 3-A(12)(d). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

I. Is the study on EPA's website	? Yes NoX
1a. If not on the EPA we	ebsite, please attach a copy.
2. Author(s) or researcher(s)	M.K. Hausbeck
	B.D. Cortright
. Publication and Date of Public	ation Research in progress
. Location of research study	Michigan, USA
	, Iodomethane, Composted chicken manure
i. Was crop yield measured in th	e study? Yes NoX
S. Was crop yield measured in th	e study? Yes No X  the alternative in controlling pests in the study.
i. Was crop yield measured in the control of the co	e study?  Yes NoX  the alternative in controlling pests in the study.  yet.  study apply to your situation. Would you expect similar results? Are there

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# Worksheet 3-A/13) Alternatives - Technical Feasibility of Alterna

orksheet or A(10). Alternatives - reclinical reasibility of Alternatives to Methyl Bromide
In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.
For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Grafting, Resistant Rootstock,	Study: UNEP 1995, UNEP 1998, UNEP 2000, B-36, B-83,
Plant Breeding	A-76, B-46, B-94, D-91, D-105, D-109, B-47, B-281
Section I. Initial Screening on Technic	cal Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the	use of this alternative on you	ır site?
1a. Full use permitted	X	62 \$10 \$25
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		

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# Worksheet 3-A(13). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

	a. If not on the EPA website, please attach a copy. s) or researcher(s)	
2. Autho		2.5
2. Autho	s) or researcher(s)	_
3. Public	tion and Date of Publication	<del></del> _
4. Locati	n of research study	
E Name	f alternative(s) in study. If more than one alternative, list the ones you wish to discuss.	
	Resistant Rootstock, Plant Breeding	
Oraitin		
6. Was c	p yield measured in the study? Yes No	
7. Descr	e the effectiveness of the alternative in controlling pests in the study.	
	••	
-		
other	s how the results of the study apply to your situation. Would you expect similar results? Are there ctors that would affect your adoption of this tool?	
The re	ults of the study do not readily apply to cucurbit production in Michigan. The study focuses on fruit trees	
grapes	for control of nematodes and soil-borne pathogens. Grafting cucurbits onto resistant rootstock will not	
solve	e problem of fruit rot of cucurbits. Currently, resistance has not been identified. (See information under	
Alterna	ive #15.)	

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## Worksheet 3-A(14). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used
—successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible
alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Organic Production	Study: UNEP 1998, B-91
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## Section I. Initial Screening on Technical Feasibility of Alternatives

1. /	Are i	there any	location-spe	ecific restrictio	ns that inhibit	t the use of th	is alternative on	your site?
------	-------	-----------	--------------	-------------------	-----------------	-----------------	-------------------	------------

1a.	Full use permitted	X	
1b.	Township caps		975 1 12 20 20 20 20 20 20 20 20 20 20 20 20 20
1c.	Alternative not acceptable in consuming country		
1d.	Other (Please describe)		192 na. 1944

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# Worksheet 3-A(14). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide No \_\_\_\_ Yes X 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Organic Production No \_\_\_\_ 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Michigan cucurbit growers would/not expect the same results since they are not able to use solarization to reduce initial inoculum of Phytophthora capsici in the soil. The other aspects of organic production are currently being used by Michigan growers but do not provide adequate protection.

# Worksheet 3-A(14)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Organic Production	Study: Tomatoes and cucurbits in Egypt: Certified organic
	methods
Section I. Initial Screening o	on Technical Feasibility of Alternatives
	•

<ol> <li>Are there any location-specific restrictions that inhibit the</li> </ol>	ne use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	

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# Worksheet 3-A(14)(b). Alternatives - Technical Feasibility of Alternatives to Methyl **Bromide** Section II. Existing Research Studies on Alternatives to Methyl Bromide Yes X 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Organic Production 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study do not apply to the growing situation in Michigan because Egypt is using greenhouses and modified greenhouses, which would have no impact on the Phytophthora capsici problem. Similarly, solarization, which was used in the Egypt study, would not be effective in a cooler Michigan climate.

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## Worksheet 3-A(14)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

## BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Organic Production	Study: Alternatives for methyl bromide on cucurbits, 2	2002.
· · · · · · · · · · · · · · · · · · ·	otady. Alternatives for incliny bronning on cucurbits, 2	ZUUZ.

# Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the	use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	·
	·

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#### Worksheet 3-A(14)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

#### Section II. Existing Research Studies on Alternatives to Methyl Bromide No X Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. M.K. Hausbeck 2. Author(s) or researcher(s) B.D. Cortright Research in progress 3. Publication and Date of Publication Michigan, USA 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Telone C-35, Chloropicrin 100%, Iodomethane, Composted chicken manure 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Fields have not been harvested yet. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable, since the research was conducted in Michigan, USA.

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#### Worksheet 3-A(15). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Resistant Cultivars	Study: UNEP 1998, B-83, B-282

#### Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restrictions that inhibit the use of this alterna	itive on	your site
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1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
-		

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ID#	

#### Worksheet 3-A(15). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

### Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? No \_\_\_\_\_ 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Resistant Cultivars 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The comments in the reports B-83, B-282 do not state that genetic resistance to Phytophthora capsici exists. Our data suggest that genetic resistance has not been identified to this soil-borne pathogen.

OMB Control # 2060-0482

#### Worksheet 3-A(15)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Resistant Cultivars	Study:	Screening curcurbits	for genetic resistar	nce to fruit
		rot in pickles, 1998-2	001	

#### Section I. Initial Screening on Technical Feasibility of Alternatives

11616	any location-specific restrictions that milibit the	use of this afternative	on your site?	
1 <i>a</i>	a. Full use permitted	X		
1b	o. Township caps			
10	c. Alternative not acceptable in consuming country			
1d	l. Other (Please describe)			
			71106	

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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ID#	

### Worksheet 3-A(15)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

#### Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) M.K. Hausbeck R. Hammerschmidt 3. Publication and Date of Publication Pickle Seed Research Fund reports, 1999, 2000, 2001 4. Location of research study Michigan 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Resistant Cultivars 6. Was crop yield measured in the study? Disease resistance was measured. 7. Describe the effectiveness of the alternative in controlling pests in the study. Resistant varieties were not identified. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable to Michigan, since it was conducted locally. Results indicate that currently genetic resistance to *Phytophthora capsici* is not available.

OMB Control # 2060-0482

#### Screening Cucurbits for Genetic Resistance to Fruit Rot in Pickles, 1998-1999

Submitted by: M. Hausbeck R. Hammerschmidt

Characteristics of the challenge inoculum: Four field isolates of *P. capsici* exhibiting diversity for mating type, sensitivity to mefenoxam, host type, and origin were selected as representative of the diversity in Michigan cucurbit production fields. These include: 1) OP97, isolated from pickling cucumber fruit in northwestern Michigan in 1997, A1 mating type, and fully sensitive to mefenoxam; 2) SP98, isolated from pumpkin fruit in southwestern Michigan in 1998, A2 mating type, and fully sensitive to mefenoxam; 3) SFF3, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A2 mating type, and intermediately sensitive to mefenoxam; 4) SF3, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A1 mating type, and intermediately sensitive to mefenoxam; 5) a control was included that consisted of an agar plug only with no pathogen present.

Mating type was determined by mating each isolate to known A1 and A2 isolates on unclarified V8 agar (UCV8) plates and scoring for the presence or absence of oospores after a three to five day incubation period. Mefenoxam sensitivity was determined in vitro by placing a 0.7 mm plug of actively expanding mycelium onto the center of  $100 \times 15 \text{ mm}$  UCV8 plates amended with 0 and 100 ppm mefenoxam. Plates were incubated at 23 to  $25^{\circ}\text{C}$  for three days and colony diameter measured. Percent growth on the amended plates was determined relative to the unamended control. Percent growth <30% of the control is designated as sensitive, between 30 to 90% as intermediate sensitivity, and >90% as fully insensitive.

Fruit preparation and inoculation: Cucumbers were grown in fields with a negative history for *P. capsici* infection according to standard practices. Mature fruit were harvested weekly, sorted according to size, and stored in a cold room at 4°C until a chamber experiment could be initiated (generally three to six days). Fruit were subjected to a 5 minute immersion in a 5% commercial bleach solution and gently washed, then rinsed in distilled water. Fruit were allowed to dry under ambient conditions. Dry fruit were labeled with a numerical code on both ends with a permanent marker. A 0.7 mm plug of actively expanding mycelium or plain UCV8 agar was placed at the center of unwounded fruit.

**Experimental design:** Two incubation chambers were constructed adjacently in a single room. Incubation chambers consisted of a ten foot diameter plastic wading pool with a polyethylene cap. Temperature and relative humidity was measured with a portable HOBO sensor. Relative humidity was uniformly 100% and temperature varied between 21 and 23°C diurnally.

A completely randomized design determined the layout of inoculated fruit in individual chambers. Fruit were given a number assignment indicating the inoculum and cucumber variety.

A random number generator was used to construct a linear array of the number set used to code individual fruit. Moist cheesecloth was placed onto the floor of the incubation chamber prior to laying out the inoculated cucumbers according to this random number sequence. Fruit were incubated for three to six days and scored for lesion diameter, sporulation intensity, and the diameter of sporulation.

Each of the five treatment/host combinations was replicated either two or four times within a single experiment and each treatment/host combination was represented in at least two experiments. Initially there were four replications of each treatment/host combination, but due to a lack of space as the growing season progressed this was reduced to only two replications per experimental run. When possible, identical experiments were replicated.

Number of cucumber cultivars screened for resistance to *Phytophthora capsici* in 1999.

Cucu <u>m</u> ber type	Commercial varieties	Numbered varieties	Plant introduction varieties	Total number screened
Pickling	22	8	37	67
Slicing	6	5		11
Total	28	13	37	78

Further analysis on this year's data will be conducted to choose lines that will be evaluated in next year's screen. While all lines appeared to be susceptible, we are interested in pursuing those lines where lesion diameter and sporulation density was reduced. See Appendix II for tables.

Incubated 5 days

Variety	P. capsi	ci OP97	P. capsi	ici SP98	P. capsici SFF3		P. capsici SF3	
	Lesion diam. (cm) (ave) <sup>I</sup>	Sporulation density <sup>2</sup> (ave)	Lesion diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. Sporulation (cm) (ave) density (ave)		Lesion diam. (cm) (ave)	Sporulation density (ave)
ACX 18, Abbott & Cobb	8.2	1.8	8.4	1.0	8.7	1.5	8.2	0.5
ACX 5001, Abbott & Cobb	8.6	1.6	8.9	1.4	8.8	1.4	8.3	0.6
ACX 5002, Abbott & Cobb	8.6	0.6	8.7	1.8	8.6	2.0	8.5	0.3
Dasher II, Petoseed	8.4	0.9	8.6	1.5	8.8	1.4	8.3	0.1
General Lee, Harris Moran Seed	8.5	1.0	8.2	1.3	8.5	1.4	8.3	0.0
Greensleeves, Harris Moran Seed	8.2	2.1	8.33	$0.8^3$	$8.8^{3}$	1.33	8.6³	1.0³
Panther, Sun Seeds	9.0	1.3	8.7³	1.8	8.6	1.9	8.34	1.4⁴
Speedway, Petoseed	8.5	1.8	8.4	2.6	8.9	1.8	8.2	0.8
SRQS 2387, Sun Seeds	8.2	0.6	8.5	1.6	8.8	1.1	8.8	0.8
SRQS 2389, Sun Seeds	8.9	1.9	8.64	1.54	9.0	1.3	8.1	0.1
Ultra Pak, Stokes	8.8	0.8	8.83	2.13	8.6	1.4	8.5	0.1
Vlaspick (pickling)	8.4	2.9	8.5	2.9	8.7	2.9	8.33	2.5³

verage of two replications (four fruits per replicate).

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

hree fruits averaged in replicate 1.

wo fruits averaged in replicate 1.

Experiments 1 and 2, incubated 3 days

<sup>7</sup> ariety	P. capsici OP97		P. capsici SP98			P. capsici SFF3			P. capsici SF3			
	Lesion diam. (cm) (ave) <sup>1</sup>	Sporulation diam. (cm) (ave)	Sporulation density <sup>2</sup> (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm) (ave)	Sporulation density (ave)
Dividend, Rogers	4.6	2.8	1.0	4.3	2.5	1.3	4.4	2.8	1.0	4.5	3.5	1.5
ortune, Rogers	4.6	4.1	1.8	4.9	4.5	2.0	4.5	4.1	1.5	4.4	3.7	2.0
Golden Dawn II, Rogers	4.9	3.7	1.5	4.3	3.0	1.5	4.2	3.0	1.0	4.5	3.6	2.3
Aultipak, Harris Moran Seed	4.3	4.1	1.8	4.0	3.4	2.0	4.3	3.8	2.0	4.3	4.0	2.3
evenue, Rogers	4.7	3.6	1.8	3.9	3.2	1.8	4.9	3.7	2.0	4.5	3.5	1.5
SQ 496-VP, Rogers	5.2	4.1	2.0	.5.0	3.8	1.8	5.2	4.0	2.5	5.8	4.7	2.3
SQ 7703-VP, Rogers	4.54	3.74	2.34	5.2	3.3	1.3	4.6	3.3	1.8	4.94	3.84	$2.0^{4}$
SQ 8057, Rogers	4.8	3.9	1.5	5.0	3.8	2.0	4.7	3.4	1.5	5.5	3.9	2.0
SQ 8058, Rogers	4.3	3.1	1.5	4.8	3.7	1.8	5.6	4.1	1.8	4.5³	$3.5^{3}$	1.53
SQ 8067, Rogers	5.4	4.3	2.0	5.1	4.1	2.3	5.6	4.2	2.0	5.1	3.5	1.8
pineless Beauty, Rogers	4.7	3.5	2.0	4.9	3.0	1.5	5.5	3.9	1.8	5.1	4.0	1.8
igress, Harris Moran Seed	5.5	4.4	2.3	4.5	4.0	2.0	5.1	3.6	1.3	5.2	3.8	1.8
ucchini Elite, Harris Moran Seed	4.5	3.6	2.5	4.8	3.7	1.8	4.9	3.6	2.0	5.0	4.0	2.3

verage of two replications (two fruits per replicate).

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit in replicate 2.

ne fruit in replicate 1.

### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici SQUASH Experiment 3, incubated 4 days

Variety (two fruits per		P. capsici OP9	07		P. capsici SP9	8 <u>aoaica 4 days</u>	:	P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Cougar, Harris Moran Seed	4.0	3.4	2.5	4.5	3.7	2.0	4.1	3.0	1.5	4.3	3.4	2.0
General Patton, Asgrow	4.2	3.5	2.0	4.4	4.0	2.5	4.6	3.3	2.0	3.9	3.4	2.5
Golden Rod, Harris Moran Seed	4.5	3.7	2.0	4.7	0.0	0.0	4.2	0.0	0.0	4.2	0.0	0.0
HMX 8714, Harris Moran Seed	4.7	2.6	1.5	4.0	1.6	1.0	4.1	0.0	0.0	4.7	2.8	2.0
HMX 8714, Harris Moran	3.3	2.2	2.0	4.3	1.6	1.5	3.9	0.0	0.0	4.4	1.4	1.0
IMX 8727, Harris Moran	3.5	1.3	0.5	4.0	2.7	1.0	4.1	1.3	0.5	3.4	1.0	0.5
HMX 9705, Harris Moran Seed	2.8	0.7	0.5	3.0	0.0	0.0	2.3 <sup>2</sup>	0.02	$0.0^{2}$	4.0	1.3	0.5
IMX 9706, Harris Moran eed	3.1	1.2	1.0	3.5	2.9	1.5	2.3	1.0	0.5	3.2	1.3	1.0
iberator II, Asgrow	5.3	3.9	1.5	, 4.4	3.0	1.5	4.3	. 2.5	1.5	4.9	3.6	2.0
fedallion, Abbott & Cobb	4.0	3.5	1.5	4.6	3.7	1.5	4.4	2.8	1.0	4.2	3.2	2.0
evenue, Rogers	3.3	0.0	0.0	3.0	1.6	1.0	3.1	0.0	0.0	1.3		2.0
SXP 709, Harris Moran Seed	3.7	2.8	2.0	3.6	2.7	1.5	3.7	3.0	2.5		0.0	0.0
SXP 787, Harris Moran Seed	5.2	3.1	1.5	5.0	3.0	2.0	3.2	0.0		3.8	3.8	2.0
SXP 788, Harris Moran Seed	4.7	2.5	1.5	4.9	3.0	1.5	4.1		0.0	4.3	1.8	1.5
	······································		···· <u>·</u>	<del></del>		1.5	7.1	0.0	0.0	4.7	1.9	1.0

#### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH** Experiment 3, incubated 4 days

continued

SXP 789, Harris Moran	4.1	3.1	1.5	4.8	2.8	2.5	3.5	0.6	0.5	3.0	2.0	1.5
SXP 798, Harris Moran	3.5	2.4	1.0	3.6	2.3	1.0	3.6	1.6	1.0	3.5	2.3	1.5
ıperpik, Harris Moran ed	3.7	3.4	3.0	4.4	3.1	3.0	4.2	3.2	1.5	4.1	3.1	2.5
ıperset, Harris Moran eed	4.0	3.1	1.5	$3.8^{2}$	2.82	$1.0^{2}$	4.6	3.0	1.0	3.5	2.7	2.0

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. e fruit per replicate.

Experiment 4, incubated 3 days

ariety (two fruits per		P. capsici OP9	<i>i</i>		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (avc.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
ougar, Harris Moran Seed	4.9	4.1	1.5	5.0	3.2	2.0	4.9	3.3	1.5	4.2	2.3	1.5
eneral Patton, Asgrow	4.9	4.4	2.5	5.0	5.0	2.0	4.6	4.6	1.0	5.5	5.2	1.5
olden Rod, Harris Moran	3.6	3.4	1.0	3.7²	2.72	$2.0^{2}$	4.6	3.4	1.0	3	3	3
MX 8714, Harris Moran eed	5.2	3.5	2.5	4.7	3.4	2.0	5.4	1.5	1.0	5.0	3.4	2.0
MX 8714, Harris Moran eed	3.6²	2.5 <sup>2</sup>	$2.0^{2}$	5.4	3.4	2.0	5.6	3.1	2.0	5.4 <sup>2</sup>	3.7 <sup>2</sup>	$3.0^{2}$
MX 8727, Harris Moran	4.3	4.3	1.0	4.7 <sup>2</sup>	$3.9^{2}$	2.0 <sup>2</sup>	4.5²	0.02	$0.0^{2}$	4.6 <sup>2</sup>	2.82	1.0²
MX 9705, Harris Moran eed	4.2²	4.22	$1.0^{2}$	3.32	3.3 <sup>2</sup>	$2.0^2$	3	3	3	3	3	3
MX 9706, Harris Moran eed	4.2	3.8	1.5	4.7	3.8	1.5	4.6	3.4	1.5	4.7	4.0	1.5
iberator II, Asgrow	5.3	4.9	2.0	<sup>'</sup> '5.7	5.5	1.5\	5.8	5.3	2.0	5.5	5.5	2.0
fedallion, Abbott & Cobb	4.2	4.2	2.0	5.0	4.6	2.0	5.7	5.2	3.0	4.7	4.0	3.0
evenue, Rogers	4.1	3.1	1.0	5.7	4.8	1.0	5.1	3.7	1.5	5.9	4.6	2.0
SXP 709, Harris Moran Seed	4.5	4.1	3.0	4.8	4.5	2.0	6.0	5.3	2.5	5.7	5.4	2.5
SXP 787, Harris Moran Seed	5.3	3.1	1.5	5.4	2.3	1.0	5.2	3.2	1.5	5.3	4.1	2.0
SXP 788, Harris Moran Seed	5.0	2.9	1.0	3.9	3.1	2.5	5.0	2.9	1.5	4.7	4.7	1.5

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#### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH** Experiment 4, incubated 3 days

continued

SXP 789, Harris Moran	4.9	3.6	3.0	-5.4	3.7	2.0	4.5	3.4	2.5	5.0	2.7	1.5
SXP 798, Harris Moran Seed	3.9	3.1	2.0	4.6	4.1	1.0	4.9	3.1	2.0	4.1	3.0	1.0
uperpik, Harris Moran Seed.	4.7	3.6	2.0	4.9	3.2	1.5	4.5	3.5	1.0	4.8	4.6	2.0
uperset, Harris Moran Seed .	5.7	5.4	2.5	5.5	5.5	2.5	5.7	3.4	1.5	5.7	4.4	1.5

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit per replicate.

ruits were too contaminated to evaluate.

Experiment 5, incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion dïam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
CX 34, Abbott & Cobb	5.7	4.6	2.5	6.1²	3.5 <sup>2</sup>	$2.0^{2}$	5.4	4.2	2.0	5.7	4.4	3.0
vidend VIP, Siegers Seed	5.1	4.2	3.0	5.9	5.0	2.0	6.4	4.6	2.0	5.7	4.5	3.0
olden Dawn	4.6	3.6	1.5	4.5	3.8	1.5	4.9	3.4	1.0	4.5	3.4	2.0
evenue, Siegers Seed	5.9	4.7	2.0	5.7	4.4	2.0	5.5	4.1	2.0	5.3	4.5	2.5
KS 9732, Sun Seeds	5.8	4.5	3.0	5.5	3.8	2.0	5.6	3.8	2.0	5.3	4.3	2.5
RHT 1777	4.2	3.6	2.0	4.1	3.0	1.0	4.6	4.1	2.0	4.8	3.8	2.5
icchini Elite F1, Harris oran Seed	5.2	3.9	2.0	5.5	4.5	2.0	4.7	3.2	2.0	4.7	3.5	2.0

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

e fruit per replicate.

Pickling cucumber Experiment 4, incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9			P. capsici SFF	3		P. capsici SF3	<u> </u>
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) '(ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
.CX 34, Abbott & Cobb	4.9	3.5	2.0	4.8	3.3	2.5	4.5 <sup>2</sup>	3.0 <sup>2</sup>	2.0 <sup>2</sup>	5.0	3.3	2.0
Dividend VIP, Siegers Seed	3.6	0.6	0.5	5.0	3.9	2.0	4.0	1.5	0.5	4.6	3.0	2.0
evenue, Siegers Seed	4.5	3.2	2.0	4.8	3.4	2.0	5.2	3.7	2.0	4.6	2.8	1.5
easons, Abbott & Cobb	3.6	1.9	1.5	5.2	1.8	0.5	5.9	3.6	2.0	5.4	3.8	1.5
eneca Prolific, Siegers Seed.	4.9	4.1	2.0	5.4	4.5	1.5	5.3	4.1	2.0	4.6	1.8	0.5
XS 9732, Sun Seeds	5.6	4.2	3.0	5.4	3.9	1.5	5.0	3.7	2.0	4.9	3.2	2.5
ucchini Elite F1, Harris Ioran Seed	5.2	3.3	1.0	4.2 •	3.3	1.5	5.3	3.6	2.0	4.0	1.8	1.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. ne fruit per replicate.

Commence of the Mindelphia.

Rerun of pickling cucumber Experiment 4, incubated 3 days

ariety (two fruits per eplicate)		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
opinoutoj	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
ACX 34, Abbott & Cobb	4.6	3.8	1.5	5.1	3.8	1.0	4.8	3.8	1.5	2	2	-2
Dividend VIP, Siegers Seed	4.5	3.3	2.5	4.7	4.1	1.0	4.8	3.7	1.5	4.2	1.9	0.5
Levenue, Siegers Seed	3.8	1.7	1.0	4.4	1.7	0.5	4.8	1.7	1.0	3.6	1.4	1.0
easons, Abbott & Cobb	4.0	1.8	1.0	5.1	4.3	1.0	, 4.5	3.9	1.0	3.5	1.6	0.5
eneca Prolific, Siegers Seed .	4.8	3.9	3.0	4.7	4.2	2.5	4.7	3.8	2.5	3.8	3.3	2.0
XS 9732, Sun Seeds	4.9	3.7	2.5	4.4	3.3	1.0	2.5	1.3	0.5	4.4	3.4	1.5
ucchini Elite F1, Harris Ioran Seed	4.6	3.2	1.5	4.2	2.7	1.0	4.0	1.5	0.5	4.5	2.0	1.5

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. ruits were too contaminated to evaluate.

Pickling cucumber Experiment 6, incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
3367, USDA	5.4	3.9	1.0	3.8	0.0	0.0	3.9	0.0	0.0	5.6	1.0	0.5

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to *Phytophthora capsici* SQUASH

Pickling cucumber Experiment 7 (redo's), incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8	<u> </u>	P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam (cm) (ave.)	Sporulation density (ave.)
3367, USDA	3.5	2.6	1.5	4.1	2.2	1.	5.3	3.2	1.0	4.3	2.5	1.0
oldfinger	5.0²	$3.3^{2}$	$1.0^{2}$	3	3	3	4.2 <sup>2</sup>	$3.3^{2}$	$1.0^{2}$	3.5	2.3	1.0
olden Dawn III	3	3	3	3	3	3	4.2²	4.22	1.02	3.1 <sup>2</sup>	$2.0^{2}$	1.0 <sup>2</sup>
RHT 1777	5.5	4.9	2.0	4.6	3.7	2.0	5.1	4.3	1.5	4.5	3.3	1.5

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit per replicate.

uits were too contaminated to evaluate.

Experiment 1, incubated 6 days

riety (four fruits per		P. capsici OP	97		P. capsici SP9	8		P. capsici SFF	3		P. capsici SI	73
licate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	 Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
lypso, Atlas Seeds	9.1	8.6	3.0	8.34	5.24	3.04	9.6	8.7	3.0	8.7³	7.2³	$3.0^{3}$
rolina, Atlas Seeds	9.8	8.3	3.0	9.5 <sup>4</sup>	7.54	$3.0^{4}$	10.1	8.1	3.0	8.7	7.2	2.8
oss Country F1, Harris oran Seed	9.0	8.6	3.0	8.03	6.1 <sup>3</sup>	3.03	9.6	8.8	3.0	8.83	6.23	2.73
scover M Hybrid, Asgrow .	9.7	8.2	3.0	10.1	6.8	3.0	10.6	7.7	3.0	11.3	7.8	3.0
1914 183491, Seminis	9.1	8.6	3.0	7.7²	4.82	$2.5^{2}$	9.5	8.1	3.0	9.4	8.9	3.0
cel M, Asgrow	9.5	7.8	3.0	8.64	5.44	$2.0^{4}$	11.6	8.1	3.0	8.7³	$7.0^{3}$	3.0 <sup>3</sup>
ncipak, Asgrow	10.1	9.1	3.0	8.9	6.3	3.0	10.9	'9.1	3.0	10.3	8.4	3.0
4X 5020 F1, Harris Moran	9.93	$8.4^{3}$	$3.0^{3}$	7.5²	4.92	2.52	9.8	7.6	3.0	8.4	6.1	2.8
AX 3469 F1, Harris Moran	9.5	7.1	3.0•	7.62	4.02	1.52	9.8	8.2	3.0	9.03	5.9 <sup>3</sup>	2.33
AX 8460 F1, Harris Moran	10.4	8.2	3.0	9.3	6.4	2.8	10.7	9.0	3.0	9.7	7.6	3.0
AX 8461 F1, Harris Moran	10.9	8.9	3.0	9.2	6.5	3.0	10.4	9.4	3.0	9.5	8.0	3.0
fayette Classic, Sun Seeds .	10.1	8.4	3.0	7.4	4.8	2.8	9.4	8.6	3.0	10.7	8.0	3.0
209064, USDA	10.1	8.9	3.0	7.6²	4.82	$3.0^{2}$	9.8	8.3	3.0	9.4	8.7	3.0
426169, USDA	10.1	7.3	3.0	8.5 <sup>4</sup>	6.24	3.04	10.0	7.6	3.0	10.0	6.6	2.8
466922, USDA	9.6	7.7	3.0	8.8	6.4	. 3.0	9.6	. 8.3	3.0	10.3	8.0	: : : 3: <b>0</b> : : :
oneer, Atlas Seeds	9.4	8.3	3.0	7.6²	6.12	$3.0^{2}$	9.9	8.8	3.0	9.8	6.5	2.5

Experiment 1, incubated 6 days Continued

riety (four fruits per dicate)		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	73
	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
gal F1, Harris Moran Seed .	10.4	8.4	3.0	9.2³	7.43	$3.0^{3}$	10.8	8.6	3.0	10.6	8.4	3.0
yal Fl, Harris Moran Seed .	9.6	8.8	3.0	8.9³	$7.5^{3}$	$3.0^{3}$	9.3	8.6	3.0	10.2	8.4	3.0
QP 2391, Sun Seeds	10.4	9.0	3.0	<b></b> 5	5	5	9.2	8.5	3.0	8.7	7.8	3.0
ıllion 193782, Seminis Seed	9.9	8.5	3.0	10.7	8.0	3.0	10.3	9.5	3.0	11.4	9.5	3.0
mter, Atlas Seeds	9.9	7.8	3.0	9.1 <sup>2</sup>	$6.2^{2}$	$3.0^{2}$	1,1.0	8.0	3.0	9.7	7.2	3.0
mor Hybrid, Asgrow	10.7	9.5	3.0	10.4	8.7	3.0	10.4	9.2	3.0	10.0	9.1	3.0
ansamerica F1, Harris Moran	9.7	7.7	3.0	8.5²	$7.0^{2}$	$3.0^{2}$	10.8	8.4	3.0	9.2	8.8	3.0
ctoria, Sun Seeds	10.3	8.1	3.0	9.4	9.0	3.0	10.33	9.2³	$3.0^{3}$	11.4	9.8	3.0
aspik VGA733, Seminis	10.2	7.5	3.0	9.3	5.7	2.5	10.7	7.4	3.0	9.1	6.5	2.5
asset, Asgrow	8.7	8.1	3.0	8.7	6.7	3.0	9.9	8.7	3.0	9.2	8.2	3.0
asspear Hybrid, Asgrow	9.6	8.4	3.0	8.8 <sup>3</sup>	$6.6^{3}$	$3.0^{3}$	10.1	,9.0	3.0	9.1	8.2	3.0
asstar B, Asgrow	9.2	7.6	3.0	6.9²	3.9 <sup>2</sup>	1.5 <sup>2</sup>	10.4	8.4	3.0	9.1	7.0	3.0
isconsin, Atlas Seeds	9.6	7.4	3.0	$9.0^{2}$	8.2 <sup>2</sup>	3.0 <sup>2</sup>	10.3	8.2	3.0	10.9	8.4	3.0

rulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

fruits were too contaminated to evaluate.

o fruits per replicate.

ee fruits per replicate.

e fruit per replicate.

Experiment 2, incubated 3 days

ariety (four fruits per		P. capsici OP9	97	٠,	P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
arolina, Atlas Seeds	3.9 <sup>3</sup>	2.83	2.03	4.2	2.6	1.8	4.6	1.6	1.0	3.8	2.3	1.5
ross Country F1, Harris Ioran Seed	4.9³	3.33	2.03	3.5	2.4	1.33	4.8	2.5	1.3	4.3	3.0	1.5
iscover M Hybrid, Asgrow .	4.7	1.6	1.0	4.2	1.8	1.0	5.0²	$3.0^{2}$	1.52	5.1	1.8	1.3
X 1911 155633, Seminis	4.5	2.9	2.0	4.1	2.6	2.0	4.3²	3.4 <sup>2</sup>	$2.0^{2}$	4.2³	$2.8^{3}$	1.33
X 1914 183491, Seminis	4.7³	$3.4^{3}$	$2.0^{3}$	4.5³	3.33	$2.0^{3}$	3.8	3.1	2.0	4.5 <sup>2</sup>	3.3 <sup>2</sup>	2.5 <sup>2</sup>
xcel M, Asgrow	4.2	2.7	1.5	4.8	3.0	1.8	5.0 <sup>3</sup>	3.13	$1.7^{3}$	4.3³	$2.9^{3}$	1.73
MX 3469 F1, Harris Moran	4.5³	3.43	$2.0^{3}$	3.72	3.12	2.02	4.44	2.84	1.04	6.0²	4.32	3.0 <sup>2</sup>
MX 8460 F1, Harris Moran	5.2	3.4	2.0	4.9	2.8	1.5	4.6 <sup>2</sup>	3.5 <sup>2</sup>	$2.0^{2}$	4.8	2.8	1.8
MX 8461 F1, Harris Moran	4.1	3.0	1.8	4.4	2.9	1.73	4.7³	3.13	1.33	4.5³	3.13	$2.0^{3}$
ackson, Sun Seeds	4.9	3.9	2.0	5.2	2.2	1.3	4.9	3.4	2.0	5.3³	4.43	2.73
afayette Classic, Sun Seeds .	5.0	4.1	2.3	4.4 <sup>3</sup>	$2.8^{3}$	$2.0^{3}$	5.8³	$2.7^{3}$	$1.7^{3}$	$4.6^{3}$	$3.3^{3}$	$2.0^{3}$
I 466922, USDA	$5.0^{2}$	$3.4^{2}$	1.5 <sup>2</sup>	4.1 <sup>2</sup>	2.2 <sup>2</sup>	$1.5^{2}$	4.8²	3.62	$2.0^{2}$	$3.9^{3}$	$1.9^{3}$	1.33
egal F1, Harris Moran Seed.	3.9³	$3.4^{3}$	$2.0^{3}$	4.8 <sup>3</sup>	$3.3^{3}$	1.73	5.2³	$3.2^{3}$	$1.3^{3}$	4.2³	$2.7^{3}$	$1.7^{3}$
oyal F, Harris Moran Seedl .	4.1 <sup>3</sup>	$2.8^{3}$	$1.0^{3}$	4.5³	3.13	$2.0^{3}$	4.53	$3.0^{3}$	1.73	$3.6^{2}$	2.12	1.5 <sup>2</sup>
tallion 193782, Seminis	4.6²	3.1 <sup>2</sup>	$2.0^{2}$	2.54	2.54	2.04	3.3³	2.83	1.73	4.6	3.5	2.0
amor Hybrid, Asgrow	4.2	2.5	1.5	4.1 <sup>2</sup>	2.82	1.5 <sup>2</sup>	4.7³	$3.4^{3}$	$2.0^{3}$	3.2 <sup>2</sup>	2.4 <sup>2</sup>	$1.0^{2}$
lasstar B, Asgrow	4.5	3.1	2.0	3.5	1.8	1.0	4.9	2.2	1.0	4.5	3.0	1.8

Experiment 2, incubated 3 days
Continued

ariety (four fruits per		P. capsici OP9	7	P. capsici SP98				P. capsici SFF.	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density! (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
isconsin, Atlas Seeds	5.4	2.1	1.0	4.4	3.0	1.8	4.9²	2.9²	1.52	5.2	3.1	1.8

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

o fruits per replicate.

ree fruits per replicate.

e fruit per replicate.

Experiment 3, incubated 4 days

'ariety (two fruits per eplicate)	•••	P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
apricate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Calypso, Atlas Seeds	6.0	4.1	2.0	6.3	4.5	1.5	6.9	4.9	2.5	5.9	3.7	2.0
X 1911 155633, Seminis	6.3	4.3	2.0	5.4	4.9	2.0	5.9	4.7	2.0	7.0	4.8	2.0
'ancipak, Asgrow	6.9	3.9	1.5	7.6	4.1	2.0	6.3	5.1	2.0	5.9	3.4	1.5
'MX 5020 F1, Harris Moran leed	6.6	5.0	2.5	6.8	4.9	2.0	6.6	5.1	2.5	6.1	5.3	2.5
'I 209064, USDA	6.9	4.0	1.5	6.9	5.2	2.0	6.4	4.4	2.0	6.4	4.3	1.5
יו 390241, USDA	6.1 <sup>2</sup>	6.12	$3.0^{2}$	6.1	5.6	3.0	5.5 <sup>2</sup>	6.3 <sup>2</sup>	$1.0^{2}$	5.5 ·	√. 4.9 ° ±	
'I 391570, USDA	5.9²	4.6 <sup>2</sup>	$2.0^{2}$	4.5	3.9	2.0	5.7	4.9	2.0	3.3 <sup>2</sup>	3.3 <sup>2</sup>	2.02
'I 422182, USDA	6.2	4.8	2.0	5.8	1.8	0.5	6.8	5.6	2.5	6.8 <sup>2</sup>	3.7 <sup>2</sup>	2.0 <sup>2</sup>
'I 426169, USDA	5.9	3.9	1,5	5.5	2.3	1.0	6.1	4.2	1.5	5.9	5.0	2.0
'I 426170, USDA	5.8	3.9	1.5	5.7	3.4	1.5	6.0	4.2	2.0	6.0	3.3	1.0
'I 432890, USDA	5.1	4.6	2.0	5.2	3.3	1.5	5.6²	5.6 <sup>2</sup>	$2.0^{2}$	6.4	5.0	2.0
'I 483339, USDA	5.6²	$3.8^{2}$	$2.0^{2}$	6.0	4.4	1.5	5.3	4.3	2.0	4.2	3.4	2.0
'ioneer, Atlas Seeds	6.5	4.7	4.0 .	5.9	2.5	1.5	7.7	5.2	1.0	4.7	4.7	2.0
RQS 2389, Sun Seeds	5.2	5.2	2.5	6.4	5.8	2.5	5.6	4.7	2.5	7.1	5.5	2.0
SRQP 2391, Sun Seeds	4.7 <sup>2</sup>	4.7 <sup>2</sup>	$2.0^{2}$	4.2 <sup>2</sup>	4.2 <sup>2</sup>	$2.0^{2}$	6.6	5.1	3.0	7.0	4.0	2.0
ransamerica F1, Harris Moran Seed	6.0	4.2	2.0	6.0	4.1	2.0	5.1	3.7	1.5	5.5	4.1	2.0
/laspik VGA733, Seminis	6.7	2.3	1.0	5.8	3.6	2.0	7.0	4.9	2.0	6.3	2.9	1.5
/lasset, Asgrow	5.5	5.5	3.0	5.3	4.9	2.5	5.9	5.0	2.0	6.0	4.1	2.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit per replicate.

Experiment 4, incubated 3 days

'ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
I 167223, USDA	3.7	2.8	1.5	4.8	3.0	1.0	4.3	2.5	1.0	3.3²	$0.0^{2}$	$0.0^{2}$
1 197086, USDA	3.4	2.1	1.0	3.8 <sup>2</sup>	2.72	$2.0^{2}$	4.4	2.1	1.0	3.2 <sup>2</sup>	1.62	1.0 <sup>2</sup>
PI 209069, USDA	3.5	0.0	0.0	3.4	0.0	0.0	4.9	2.5	1.5	3.7	1.0	0.5
PI 234517, USDA	5.0	0.0	0.0	4.3²	1.92	$2.0^{2}$	3	3	3	1.5 <sup>2</sup>	$1.5^{2}$	1.0 <sup>2</sup>
PI 271328, USDA	5.0 <sup>2</sup>	2.12	$1.0^{2}$	3	••³	3	3	3	3	3.3 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
PI 288238, USDA	3	3	3	5.0 <sup>2</sup>	$3.4^{2}$	$2.0^{2}$	5.3 <sup>2</sup>	3.1 <sup>2</sup>	$2.0^{2}$	4.0	2.8	1.5
PI 330628, USDA	4.3	2.1	1.5	4.0	1.2	0.5	4.6	3.0	1.5	4.4	1.5	0.5
PI 390241, USDA	3.4	1.2	0.5	3.4	2.0	1.0	4.3	2.1	1.0	3	3	3
PI 391570, USDA	3.6²	$0.0^{2}$	$0.0^{2}$	3	3	3	4.12	2.12	1.02	4.2²	$3.0^{2}$	1.0 <sup>2</sup>
PI 422182, USDA	4.1	1.9	1.0	4.6	2.3	1.0	4.5	2.7	1.0	4.0	2.2	1.0
PI 426170, USDA	4.0	1.8	1.0	3.7 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$	4.2²	$0.0^{2}$	$0.0^{2}$	2.1 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
PI 432851, USDA	3	3	<sup>3</sup>	3	·3 .	3	3	3	3	3	3	3
PI 432855, USDA	3.6 <sup>2</sup>	2.5	$2.0^{2}$	, ,3	3	-÷³ \	3	3	3	3	3	3
PI 483339, USDA	4.4	1.2	0.5	4.7 <sup>2</sup>	$2.9^{2}$	$1.0^{2}$	3	3	3	3.6 <sup>2</sup>	2.2 <sup>2</sup>	1.0 <sup>2</sup>
SRQS 2389, Sun Seeds	4.1	2.7	2.0	4.0	2.2	1.5	4.0	2.2	1.5	3.8	2.8	1.5

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit per replicate.

Il fruits were too contaminated to evaluate.

The state of the state of the

Experiment 5, incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
I 163213, USDA	4.6	2.3	1.0	4.9	2.5	1.0	5.2	2.2	1.0	4.2	2.3	1.0
I 167223, USDA	5.4	3.1	1.5	5.6	3.5	1.0	8.1	3.5	1.0	5.0	1.5	0.5
I 197086, USDA	3.1	0.5	0.5	2.6	2.6	1.0	3.4	2.2	1.5	4.0	2.6	2.0
1 197088, USDA	3.4	1.0	0.5	5.0	3.2	1.5	3.5	0.0	0.0	3.4	0.0	0.0
I 209069, USDA	5.4	3.3	1.5	4.9	3.0	1.5	4.6	2.6	1.0	4.7	2.2	1.5
i 227209, USDA	4.6	2.1	1.0	5.2	3.1	1.5	4.8	2.4	1.0	5.3	3.3	1.5
1 234517, USDA	4.8	2.9	1.5	3.9	1.3	0.5	3.9	0.9	0.5	3.8	1.4	0.5
ri 267942, USDA	. 4.6	2.9	1.5	5.2	3.2	1.0	4.8	2.1	1.0	4.2	2.2	1.0
PI 271328, USDA	3.7	0.0	0.0	5.3	1.3	0.5	3.7	1.2	0.5	4.1	0.0	0.0
PI 288238, USDA	5.4	3.2	0.5	5.7	3.7	2.0	5.6	1.7	1.5	4.9	2.7	1.0
ri 330628, USDA	4.2	2.7	2.0	4.5	2.5	1.5	4.3	1.3	0.5	2.8	0.0	0.0
PI 390244, USDA	4.6	2.6	1.5	4.9	2.8	1.0	4.7	0.0	0.0	4.4	2.1	1.0
YI 390529, USDA	5.0	1.9	1.0	4.2	1.9	1.0	5.7	3.1	2.0	3.9	2.0	1.0
1 418964, USDA	5.1	2.8	1.5	5.0	2.7	1.5	4.5	2.2	1.0	3.6	2.7	1.0
PI 432851, USDA	5.8	3.7	2.0	4.6	2.5	1.0	4.2	1.4	1.0	4.6	2.5	0.5
PI 432855, USDA	5.5	3.4	3.0	5.5	3.2	1.5	5.8	3.1	1.5	4.9	2.8	1.5
1 432865, USDA	5.7	4.5	3.0	4.8	3.7	2.0	4.4	2.6	1.0	4.5	3.0	1.5
PI 432890, USDA	4.5	2.7	1.0	4.7	1.7	1.0	6.3	1.7	1.0	5.0	2.4	1.0
Sumter, Atlas Seeds	4.7	3.4	2.0	5.0	3.2	1.5	5.5	2.4	1.0	4.9	2.8	2.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Experiment 6, incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
mes 7118, USDA	4.5	3.4	3.0	4.0	3.0	1.0	4.6	1.5	0.5	3.7	1.2	0.5
ckson, Sun Seeds	5.1	3.3	2.5	4.7	3.2	1.0	5.7	3.7	1.5	4.2	2.3	2.0
163213, USDA	4.4	1.1	0.5	5.0	1.3	0.5	4.8	0.0	0.0	4.9	0.0	0.0
197088, USDA	3.5²	$2.0^2$	1.02	4.0	1.3	0.5	3.7	0.0	0.0	4.3	0.8	0.5
(211979, USDA	5.7	2.9	2.0	5.2	1.4	0.5	4.5	0.0	0.0	4.4	0.7	0.5
[ 249562, USDA	4.1	0.0	0.0	4.3	1.0	0.5	4.9	0.8	0.5	4.6	0.0	0.0
267942, USDA	4.3	0.9	0.5	5.4	2.0	1.0	4.1	1.2	0.5	3.3	0.0	0.0
279466, USDA	5.1	2.2	1.0	5.2	1.2	0.5	4.9	1.9	1.0	3.9 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
279467, USDA	4.4	0.7	0.5	4.1	0.0	0.0	3.1	0.0	0.0	4.4	1.0	0.5
[ 27946 <b>8,</b> USDA	4.8	1.0	0.5	4.8	1.1	0.5	3.7	0.0	0.0	2.9	0.0	0.0
I 321008, USDA	5.9	0.0	0.0	5.5	1.2	0.5	5.2	0.0	0.0	4.8	` `.0.9` `.	0.5
I 390240, USDA	5.0 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$	4.7²	$0.0^{2}$	$0.0^{2}$	6.3	1.5	0.5	5.2²	1.5 <sup>2</sup>	$1.0^{2}$
1 390244, USDA	4.4	1.9	1.0	5.2	0.7	0.5	4.5	1.0	0.5	5.0	0.8	0.5
I 390246, USDA	5.4	2.1	1.0	5.6	2.4	1.0	5.6	2.7	1.0	4.3	0.0	0.0
1 390262, USDA	3.7	2.4	1.0	3.6	2.0	1.0	5.2	2.3	1.0	4.5²	$2.0^{2}$	$1.0^2$
l 390529, USDA	4.9	2.4	1.0	4.7	1.1	0.5	5.0	0.0	0.0	5.2	1.0	0.5
l 418964, USDA	5.2	2.2	1.0	6.2	2.8	1.0	4.7	2.4	1.0	3.8	0.8	0.5
432865, USDA	4.6	3.1	3.0	3.8	1.9	1.0	5.8	2.9	1.0	4.8	2.9	2.0
l 432867, USDA	5.5	2.9	1.0	5.4	2.9	1.0	5.1	1.0	0.5	4.9	0.0	0.0

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. ie fruit per replicate.

Experiment 7 (redo's), incubated 3 days

ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
plicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
.mes 7118, USDA	4.6	3.3	2.0	4.0	2.9	1.5	4.2	2.5	1.5	4.0	3.1	2.0
xcel M, Asgrow	4.2	0.0	0.0	5.6	0.0	0.0	4.5	0.0	0.0	3.6	0.0	0.0
MX 3469 F1, Harris Moran	4.8	0.6	0.5	4.5	0.0	0.0	4.4	0.0	0.0	3.2	0.0	0.0
I 211979, USDA	5.2	2.8	1.5	5.3	3.1	1.0	4.7	3.4	1.0	4.6	2.3	1.0
I 279468, USDA	5.1	4.6	1.5	4.6	3.3	2.5	4.9	3.8	2.0	4.6	3.7	2.0
I 390241, USDA	5.2	3.8	3.0	5.3	4.0	2.0	4.0	3.4	2.0	4.3	3.1	1.0
I 390246, USDA	5.0	2.9	1.0	5.1	3.0	1.0	4.7	2.4	1.0	6.1	4.4	1.5
I 422182, USDA	3.9	0.0	0.0	4.3	0.0	0.0	4.0	. 0.0	0.0	1.9	0.0	0.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

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Rerun of Experiment 4, incubated 3 days

'ariety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	<b>3</b> . % p. 3.4% p. 3.4%
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
I 167223, USDA	4.9	3.4	2.5	4.6	2.6	1.0	4.4	3.2	1.5	4.8	3.3	2.0
I 197086, USDA	4.0	2.5	2.0	3.8	1.2	0.5	4.0	2.2	1.5	3.2	0.0	0.0
I 209069, USDA	3.9	0.0	0.0	4.5	1.2	0.5	3.9	1.8	1.0	3.7	0.0	0.0
I 234517, USDA	3.3	3.3	3.0	4.2	2.9	1.0	3.8	2.9	2.0	2.8	0.0	0.0
I 271328, USDA	3.6	0.0	0.0	4.2	2.2	1.0	4.1	0.0	0.0	3.6	1.4	0.5
I 288238, USDA	5.2	3.9	1.5	4.7	3.7	2.0	4.9	3.8	3.0	4.3	3.1	2.5
I 330628, USDA	4.1	2.4	2.0	4.5	1.2	0.5	4.5	2.8	1.5	3.5	0.0	0.0
I 391570, USDA	4.6	2.8	1.0	4.5	2.3	1.0	3.3	0.0	0.0	3.9	3.0	2.0
I 426170, USDA	4.5	1.3	0.5	4.3	3.0	1.5	4.5	0.0	0.0	3.6	1.5	0.5
I 432851, USDA	4.7	2.8	2.0	4.8	3.6	3.0	4.7	1.4	0.5	4.3	1.6	0.5
I 432855, USDA	4.0	3.5	2.0	4.4	3.1	2.0	4.7	3.4	2.5	4.2	3.0	1.0
I 483339, USDA	. 4.5	2.3	1.5	4.6	3.4	1.5	3.4	1.4	1.0	4.6	1.7	0.5
RQS 2389, Sun Seeds	3.3	0.9	1.0	5.1 <sup>2</sup>	2.72	1.0 <sup>2</sup>	3.2	1.7	1.0	2.6	0.0	0.0

orulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. ne fruit per replicate.

Experiment 8, incubated 3 days

√ariety (two fruits per		P. capsici OP9	7	<u> </u>	P. capsici SP98	3		P. capsici SFF	3		P. capsici SF	3
eplicate)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)	Lesion diam. (cm) '(average)	Sporulation . diam. (cm) (average)	Sporulation density <sup>1</sup> (average)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density (average)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)
Calypso, Atlas Seeds	4.5	1.6	0.5	4.4	3.2	2.0	4.4	3.3	1.5	4.8	2.6	1.5
Carolina, Atlas Seeds	3.9	3.5	3.0	4.7	4.0	2.5	4.3	3.8	1.0	4.2	3.7	2.5
PI 249562, USDA	3	3	3	3	<b></b> 3	3	3	3	3	3	3	3
PI 279466, USDA	<sup>3</sup>	3	3	4.2 <sup>2</sup>	$3.2^{2}$	$1.0^{2}$	3	3	3	4.0 <sup>2</sup>	$2.5^{2}$	$2.0^{2}$
PI 279467, USDA	3	3	3	$3.9^{2}$	$3.9^{2}$	$2.0^{2}$	3	3	3	3	3	3
PI 358813, USDA	3	3	<sup>3</sup>	$4.6^{2}$	$2.0^{2}$	$1.0^{2}$	<sup>3</sup>	3	<sup>3</sup>	3	3	3
PI 390240, USDA	$3.9^{2}$	$3.2^{2}$	$1.0^{2}$	3	3	3	3.9²	2.5 <sup>2</sup>	$2.0^{2}$	3.5 <sup>2</sup>	2.72	$\sim 2.0^{2.5}$
PI 390241, USDA	5.1	3.8	1.5	4.8	3.6	2.5	4.2	3.1	1.0	5.4 <sup>2</sup>	$4.0^{2}$	2.0 <sup>2</sup>
PI 422182, USDA	4.0	3.0	2.0	4.9	3.8	2.0	6.0	3.9	1.0	4.5	3.4	2.0
PI 432867, USDA	3	3	3	3	3	3	4.1	3.4	1.0	3	3	3
PI 432890, USDA	4.6 <sup>2</sup>	$3.0^{2}$	$1.0^{2}$	3	3	3	5.0 <sup>2</sup>	2.2 <sup>2</sup>	$1.0^{2}$	4.7	3.1	1.5
PI 483339, USDA	4.5 <sup>2</sup>	$2.5^{2}$	2.0²	5.1 <sup>2</sup>	$3.5^{2}$	$1.0^{2}$	4.6²	$3.8^{2}$	1.0 <sup>2</sup>	2.5 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
SRQP 2391, Sun Seeds	4.1	3.7	2.5	4.0	4.0	2.0	4.6	4.1	2.0	4.0 <sup>2</sup>	4.0 <sup>2</sup>	$2.0^{2}$
Stallion 193782, Seminis	3.2 <sup>2</sup>	$3.2^{2}$	1.02	$3.4^{2}$	$3.4^{2}$	$2.0^{2}$	3	3	3	4.0 <sup>2</sup>	4.0 <sup>2</sup>	$2.0^{2}$

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

ne fruit per replicate.

<sup>.</sup>ll fruits were too contaminated to evaluate.

Run of Experiment 7 (redo's), incubated 3 days

nriety (two fruits per plicate)	P. capsici OP97			P. capsici SP98			<u> </u>	P. capsici SFF.	3		P. capsici SF.	3
epiicatė)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>(</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
72 13367	3.5	2.6	1.5	4.1	2.2	1.	5.3	3.2	1.0	4.3	2.5	1.0
Goldfinger	5.0 <sup>2</sup>	$3.3^{2}$	$1.0^{2}$				4.2²	3.3 <sup>2</sup>	1.0 <sup>2</sup>	3.5	2.3	1.0
XRHT 1777	5.5	4.9	2.0	4.6	3.7	2.0	5.1	4.3	1.5	4.5	3.3	1.5
Golden Dawn III					+=		4.2²	4.2 <sup>2</sup>	1.0 <sup>2</sup>	3.1 <sup>2</sup>	2.0 <sup>2</sup>	1.0²

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy, ne fruit per replicate.

## Screening Cucurbits for Genetic Resistance to Fruit Rot in Pickles, 1999-2000

Submitted by: M. Hausbeck R. Hammerschmidt

Characteristics of the challenge inoculum: Four field isolates of *P. capsici* exhibiting diversity for mating type, sensitivity to mefenoxam, host type, origin and AFLP fingerprint were selected as representative of the diversity in Michigan cucurbit production fields. These include: 1) OP97, isolated from pickling cucumber fruit in northwestern Michigan in 1997, A1 mating type, and fully sensitive to mefenoxam; 2) SP98, isolated from pumpkin fruit in southwestern Michigan in 1998, A2 mating type, and fully sensitive to mefenoxam; 3) 238, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A2 mating type, and intermediately sensitive to mefenoxam; 4) 236, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A1 mating type, and sensitive to mefenoxam; 5) a control was included that consisted of an agar plug only with no pathogen present.

Mating type was determined by mating each isolate to known A1 and A2 isolates on unclarified V8 agar (UCV8) plates and scoring for the presence or absence of oospores after a three to five day incubation period. Mefenoxam sensitivity was determined in vitro by placing a 0.7 mm plug of actively expanding mycelium onto the center of 100 x 15 mm UCV8 plates amended with 0 and 100 ppm mefenoxam. Plates were incubated at 23 to 25°C for three days and colony diameter measured. Percent growth on the amended plates was determined relative to the unamended control. Percent growth <30% of the control is designated as sensitive, between 30 to 90% as intermediate sensitivity, and >90% as fully insensitive.

Growing of fruit: Some cucumber cultivars grown in 1999 were selected on the basis of the last year's screening to be grown again in 2000, along with new varieties. Five seed lots of two cultivars ('Reisenschal' and 'Vlaspik') were selected on the basis of different seed treatments. Cucumbers were grown according to standard practices in fields with a negative history for *P. capsici* infection. Mature fruit were harvested weekly, sorted according to size, and stored in a cold room at 4°C until a chamber experiment could be initiated (generally three to six days).

Fruit preparation and inoculation: Fruit were subjected to a 5 minute immersion in a 5% commercial bleach solution and gently washed, then rinsed in distilled water. Fruit were allowed to dry under ambient conditions. Dry fruit were labeled with a numerical code indicating the inoculum and cucumber variety with a permanent marker. A 0.7 mm plug of actively expanding mycelium or plain UCV8 agar was placed at the center of unwounded fruit. A plastic microcentrifuge tube was placed over the agar plug and sealed to the fruit with petroleum jelly to maintain high humidity during initial infection of the fruit.

Experimental design: A completely randomized design determined the layout of inoculated

fruit. A random number generator was used to construct a linear array of the number set used to code individual fruit. Fruit were incubated for four days on a bench top at room temperature and scored for lesion diameter, sporulation density, and the diameter of sporulation.

Each of the five treatment/host combinations was replicated four times in Experiments 1 through 5. Due to a lack of fruit set of some cultivars, there were only two replications per experiment in the last two runs (Experiments 6,7). Each treatment/host combination was represented in at least two experiments.

Number of cucumber cultivars screened for resistance to Phytophthora capsici in 2000.

Cucumber type	Number screened
Pickling	21
Slicing	4
Plant introduction	32
Total	57

Further analysis on this year's data will be conducted to choose lines that will be evaluated in next year's screen. While all lines appeared to be susceptible, we are interested in pursuing those lines where lesion diameter and sporulation density was reduced. See Appendix II for tables.

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Variety (four fruits per replicate)  P=pickling variety	P	. capsici OP	97	P	capsici SP	98	P	. capsici 23	8	:	P. capsici 23	
s=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Arabian Hybrid, Asgrow <sup>P</sup>	6.7	0.5	0.3	6.7	0.5	0.3	6.9	0.6	0.8	6.6	0.5	0.3
Colt Hybrid, Asgrow <sup>P</sup>	6.7	0.0	0.0	7.0	0.6	0.5	6.8	0.0	0.0	6.9	1.1	0.8
Cyclone, Asgrow <sup>s</sup>	7.6	1.9	1.5	6.7	0.0	0.0	7.5	3.8	2.5	7.2	0.0	0.0
Discover, Seminis <sup>P</sup>	6.9	0.0	0.0	6.8	0.0	0.0	6.9	0.0	0.0	6.7	0.0	0.0
Eureka, Siegers <sup>P</sup>	6.3	1.9	1.3	6.0	0.0	0.0	6.7	2.4	1.8	6.6	1.4	1.0
Excel, Seminis <sup>P</sup>	7.1	1.8	1.3	7.0	0.6	0.5	7.0	0.0	0.0	6.7	0.0	0.0
Lafayette, Sun Seeds <sup>P</sup>	6.8	1.1	0.8	6.5	1.5	1.0	<b>7.0</b> ,	1.3	1.0	6.9	1.1	1.0
Lightning, Asgrow <sup>s</sup>	7.1	2.4	1.0	7.2	1.4	0.8	7.4	0.0	0.0	7.4	0.6	0.5
Palomino Hybrid, Asgrow <sup>P</sup>	7.3	1.6	1.3	7.4	0.5	0.3	7.0	2.0	1.5	6.4	1.1	0.8
PI 209067, USDA	6.7	3.4	1.3	6.5	2.3	1.0	6.6	1.6	1.0	6.5	1.6	0.8
PI 249561, USDA	7.1	0.0	0.0	6.7	0.0	0.0	6.5	0.0	0.0	6.8	0.0	0.0
PI 426169, USDA	7.4	1.9	1.5	6.8	0.5	0.3	7.0	2.3	1.3	6.6	1.5	0.8
Reisenschal (control), Vlasic Foods <sup>P</sup>	7.2	3.0	2.3	6.9	3.0	2.3	7.1	0.0	0.0	6.8	0.0	0.0
Stallion Hybrid, Asgrow <sup>P</sup>	6.7	0.6	0.5	7.0	0.5	0.5	7.4	0.0	0.0	7.0	0.6	0.5
Thunder, Asgrow <sup>s</sup>	7.4	1.3	0.8	7.3	0.5	0.5	7.5	0.6	0.5	7.4	0.8	0.5
Transamerica, Sun Seeds <sup>P</sup>	7.0	0.7	0.5	6.6	0.5	0.3	7.1	1.3	0.8	6.7	1.1	0.8
Vlaspik (B-1,SMP), Vlasic Foods <sup>P</sup>	6.1	2.5	2.3	6.7	0.0	0.0	7.4	0.0	0.0	6.5	1.6	1.3
Vlaspik (B-2,SMP,+), Vlasic Foods <sup>P</sup>	6.8	0.0	0.0	6.5	0.0	0.0	7.1	0.0	0.0	6.8	0.0	0.0

Variety (four fruits per replicate)	Р.	capsici OP	97	Р.	capsici SP	98	Р.	. capsici 23	8	I	P. capsici 22	36
P=pickling variety S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Vlaspik (B-3,SMP,E), Vlasic Foods <sup>P</sup>	6.7	0.0	0.0	6.5	0.0	0.0	6.9	0.0	0.0	6.6	0.0	0.0
Vlaspik (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	7.1	0.0	0.0	6,8 .	0.0	0.0	6.8	0.0	0.0	6.3	0.0	0.0
Vlaspik (control), Asgrow <sup>P</sup>	7.2	0.5	.0.3	6.7	0.0	0.0	7.3	0.5	0.3	6.6	0.0	0.0
Vlaspik+M Hybrid, Asgrow <sup>p</sup>	6.6	0.8	0.8	7.2	0.0	0.0	7.0	0.0	0.0	7.3	0.6	0.5
Vlasspear, Seminis <sup>P</sup>	6.6	2.5	2.0	7.2	2.0	1.5	7.5	0.0	0.0	6.9	0.0	0.0

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\*Rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety  S=slicing variety	P. capsici OP97			P. capsici SP98			P	. capsici 23	8	P. capsici 236		
	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)									
Arabian Hybrid, Asgrow <sup>P</sup>	6.8	0.5	0.3	6.6	0.5	0.3	6.9	0.0	0.0	6.9	0.0	0.0
Colt Hybrid, Asgrow <sup>P</sup>	6.7	0.0	0.0	7.0	0.6	0.5	7.0	0.0	0.0	7.3	0.6	0.5
Cyclone, Asgrow <sup>s</sup>	7.2	1.6	1.3	7.1	1.5	1.3	7.1	0.0	0.0	7.1	3.7	2.3
Discover, Seminis <sup>P</sup>	5.0	0.0	0.0	6.7	0.0	0.0	7.0	0.0	0.0	6.8	0.0	0.0
Eureka, Siegers <sup>P</sup>	7.1	1.8	1.5	6.5	0.5	0.5	6.9	2.5	2.3	6.9	2.4	2.3
Excel, Seminis <sup>P</sup>	6.9	2.4	2.3	7.0	1.8	1.5	7.2	0.9	8.0	6.8	0.0	0.0
Lafayette, Sun Seeds <sup>P</sup>	7.0	1.6	1.0	7.1	8.0	0.5	7.2	1.8	1.5	6.7	0.0	0.0
Lightning, Asgrow <sup>s</sup>	7.3	1.6	0.1	7.0	0.9	0.5	7.0	1.4	1.3	7.4	1.8	1.5
Palomino Hybrid, Asgrow <sup>P</sup>	7.1	0.8	0.8	6.9	0.0	0.0	6.8	1.3	0.8	6.7	0.9	0.8
PI 249561, USDA	7.1	0.6	0.5	7.2	0.9	0.8	7.1	0.0	0.0	7.0	0.0	0.0
PI 426169, USDA	7.1	1.4	1.3	6.8	0.8	0.5	7.1	1.5	1.0	6.9	1.8	1.5
Reisenschal (control), Vlasic Foods <sup>P</sup>	7.3	3,8	2.8	7:1	1.6	1.5	7.1	3.7	3.0	7.2	2.0	1.5
Stallion Hybrid, Asgrow <sup>P</sup>	7.1	0.8	0.5	6.7	0.0	、 0.0	7.0	0.0	0.0	7.2	8.0	0.8
Thunder, Asgrow <sup>s</sup>	7.3	1.0	0.8	7.1	1.8	1.0	7.4	1.0	0.5	7.3	0.0	0.0
Transamerica, Sun Seeds <sup>P</sup>	7.1	1.5	1.0	6.7	0.6	0.5	6.9	1.5	1.5	6.8	0.5	0.5
Vlaspik (B-1,SMP), Vlasic Foods <sup>P</sup>	6.8	2.5	2.3	6.4	0.8	0.8	6.3	2.5	2.3	6.7	1.0	0.8
Vlaspik (B-2,SMP,+), Vlasic Foods <sup>P</sup>	6.9	0.0	0.0	7.3	0.0	0.0	6.8	0.0	0.0	7.1	0.0	0.0
Vlaspik (B-3,SMP,E), Vlasic Foods <sup>P</sup>	6.7	0.0	0.0	6.6	0.0	0.0	6.7	0.0	0.0	7.1	0.0	0.0

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Variety (four fruits per replicate)  P=pickling variety  S=slicing variety	P. capsici OP97			P. capsici SP98			P. capsici 238			P. capsici 236		
	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporula- tion density (ave.)
Vlaspik (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	6.9	0.0	0.0	6.8	0.0	0.0	6.7	0.0	0.0	7.0	0.0	0.0
Vlaspik (control), Vlasic Foods <sup>P</sup>	7.0	1.6	1.3	7.1	0.9	0.8	7.1	2.7	2.3	7.0	1.7	1.5
Vlaspik+M Hybrid, Asgrow <sup>P</sup>	7.0	0.0	0.0	6.8	1.5	1.3	7.0	0.9	0.8	6.6	0.0	0.0
Vlasspear, Seminis <sup>P</sup>	6.9	1.3	0.8	6.9	0.0	0.0	7.2	0.0	0.0	6.7	0.0	0.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

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Variety (four fruits per replicate)  P=pickling variety	P. capsici OP97			P. capsici SP98			P. capsici 238			P. capsici 236		
	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Discover, Seminis <sup>P</sup>	6.6	0.0	• . 0.0	6.6	0.0	0.0	6.4	0.0	0.0	6.6	0.6	0.3
PI 209067, USDA	6.2	3.0	2.0	5.6	2.4	1.5	6.0	3.0	2.3	5.6	2.4	1.8
PI 209068, USDA	6.4	3.3	1.7	6.5	2.4	1.5	6.3	2.5	1.5	6.1	1.3	0.8
PI 209069, USDA	5.8	3.3	2.3	6.2	3.0	2.5	6.1	3.6	2.5	6.2	2.7	2.0
PI 211980, USDA	5.9	3.8	2.8	6.2	3.9	2.5	6.1	3.4	2.8	6.0	4.0	2.5
PI 271328, USDA	5.3	2.5	1.8	5.2	2.0	1.3	6.0	2.3	2.0	5.6	2.5	2.3
PI 330628, USDA	5.6	2.3	1.5	5.2	2.3	1.5	5.3	3.4	2.3	5.6	3.3	2.5
PI 358813, USDA	6.2	2.4	1.5	6.0	2.7	1.5	6.1	2.8	1.8	6.0	2.1	1.0
PI 358814, USDA	5.9	2.0	1.3	5.1	2.0	1.3	6.4	1.3	0.8	4.0	0.0	0.0
PI 390262, USDA	6.2	3.8	2.5	6.7	4.1	2.8	6.4	3.6	2.5	6.0	3.3	2.3
PI 390263, USDA	5.8	4.0	2.5	6.0	3.8	2.8	5.6	3.7	3.0	6.3	4.1	3.0
PI 426170, USDA	5.6	1.4	1.0	5.0	1.9	1.3	6.0	1.9	1.3	5.4	2.3	1.3
Reisenschal (B-1,SMP), Vlasic Foods <sup>P</sup>	6.1	2.9	1.8	6.6	3.6	2.5	6.4	3.2	2.0	6.6	2.8	2.3
Reisenschal (B-2,SMP,+), Vlasic Foods <sup>P</sup> .	6.5	3.3	2.3	6.2	2.4	1.5	6.4	2.3	1.3	5.6	2.9	2.0
Reisenschal (B-3,SMP,E), Vlasic Foods <sup>P</sup> .	6.4	2.8	2.0	6.1	2.6	1.8	6.3	3.0	2.3	6.4	3.1	2.8
Reisenschal (B-4,SMP,E,+), Vlasic Foods <sup>P</sup> *Sporulation density rated on a scale of 0 to 3 y	5.9	5.0	1.8	5.3	2.5	2.0	5.7	2.1	1.5	5.8	2.0	1.3

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	Р.	capsici SP	98	P	. capsici 23	8	1	P. capsici 23	36
s=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Discover, Seminis <sup>P</sup>	5.9	0.0	0.0	6.3	0.0	0.0	6.0	0.0	0.0	5.7	0.0	0.0
Meteor, Asgrow <sup>s</sup>	7.3	0.0	0.0	7.5	0.5	0.3	8.0	1.1	0.8	7.6	0.0	0.0
PI 197087, USDA	5.3	1.0	0.5	6.1	1.4	0.8	6.0	3.9	2.5	6.4	3.6	2.3
PI 197088, USDA	5.7	1.8	1.3	6.0	0.6	0.5	6.1	0.5	0.5	5.4	1.8	1.3
PI 209068, USDA	5.7	0.6	0.5	5.4	1.4	0.8	5.8	0.8	0.5	6.2	1.8	1.0
PI 209069, USDA	6.3	1.5	0.8	6.2	1.3	0.8	6.2	1.8	1.5	6.2	0.6	0.5
PI 211978, USDA	6.8	1.6	1.0	6.4	1.6	1.0	7.0	2.9	1.3	7.0	3.6	2.0
PI 211979, USDA	6.4	3.7	2.3	6.8	3.1	2.3	7.5	3.8	1.8	7.0	2.6	1.5
PI 211980, USDA	6.1	2.0	1.3	6.2	1.4	1.0	6.2	1.1	0.8	6.1	0.8	0.5
PI 271328, USDA	6.2	1.1	0.8	6.2	1.4	1.0	6.1	2.4	1.8	5.9	0.5	0.3
PI 279468, USDA	7.6	3.9	2.8	7.3	2.9	2.0	7.1	2.1	2.0	7.4	3.0	2.3
PI 330628, USDA	6.0	1.4	1.0	6.2	2.4	1.8	5.8	0.0	0.0	6.7	2.8	1.8
PI 358813, USDA	5.9	0.6	0.5	6.6	2.0	1.5	6.8	1.5	1.0	6.6	1.6	1.0
PI 358814, USDA	6.1	1.4	1.0	6.2	0.8	0.3	6.5	1.5	1.0	6.7	2.5	2.0
PI 390262, USDA	6.4	1.3	0.8	6.3	0.8	0.5	6.0	1.4	1.3	6.6	0.8	0.8
PI 390263, USDA	6.7	.0.6	0.5	6.1	2.5	1.8	6.0	0.0	0.0	6.1	0.6	0.5
PI 426170, USDA	5.9	0.6	0.5	6.0	1.1	0.8	5.6	2.2	1.3	6.0	1.3	1.0
PI 432868, USDA	7.2	3.3	2.5	7.1	2.8	2.0	7.2	3.1	2.0	6.7	2.8	1.8

Variety (four fruits per replicate)  P=pickling variety	P. capsici OP97		P	P. capsici SP98			P. capsici 238			P. capsici 236		
S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Reisenschal (B-1,SMP), Vlasic Foods <sup>P</sup>	5.7	0.5	0.3	6.1	0.0	0.0	5.7	0.6	0.3	6.4	0.0	0.0
Reisenschal (B-2,SMP,+), Vlasic Foods <sup>P</sup> .	5.9	0.0	0.0	6.2	0.0	0.0	6.0	0.0	0.0	6.1	0.0	0.0
Reisenschal (B-3,SMP,E), Vlasic Foods <sup>P</sup> .	5.9	0.0	0.0	5.6	0.0	0.0	6.0	0.0	0.0	6.1	0.4	0.3
Reisenschal (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	5.6	0.0	0.0	6.2	0.0	0.0	6.2	0.0	0.0	6.3	0.0	0.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)	Р.	<i>capsici</i> OP	97	P	. capsici SP	98	I	P. capsici 23	8	I	P. capsici 23	36
P=pickling variety S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)									
Discover, Seminis <sup>P</sup>	6.1	0.0	0.0	5.9	0.0	0.0	6.0	0.0	0.0	6.4	0.0	0.0
Meteor, Asgrow <sup>s</sup>	6.9	0.9	0.3	7.0	1.5	0.8	7.0	2.9	1.3	6.4	0.0	0.0
PI 197088, USDA	6.5	0.6	0.3	6.8	1.4	0.8	5.8	1.8	1.0	6.3	0.8	0.5
PI 211978, USDA	6.3	0.6	0.5	6.8	3.1	1.8	6.5	2.3	1.8	6.7	0.6	0.3
PI 227210, USDA	6.8	0.8	0.3	6.9	2.1	1.8	6.8	2.3	1.3	7.0	0.8	0.3
PI 279468, USDA	6.6	1.6	0.8	6.2	2.2	1.3	5.8	0.8	0.5	5.8	3.1	1.8
PI 390239, USDA	6.8	1.5	1.3	6.3	2.2	1.0	5.8	1.6	0.5	6.3	2.3	1.8
PI 432868, USDA	6.2	1.6	1.0	5.6	2.4	1.5	6.3	0.0	0.0	6.4	1.5	1.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

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1.	cupsici OF	91	P	. capsici SP	98	F	. capsici 23	8	F	P. capsici 23	16
Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm)	Sporula- tion density' (ave.)
5.4	0.0	0.0	5.2	2.1	1.0	5.1	0.0	0.0	4.9		0.0
4.8	0.0	0.0	4.8	2.8	2.5	4.1	0.0	0.0	4.7	0.0	0.0
5.6	3.6	2.5	5.3	2.3	1.0	5.5	3.0	2.0	4.9	2.3	2.0
6.1	3.1	1.5	6.1	4.0	2.5	6.1	3.1	1.5	5.9		0.0
5.6	0.0	0.0	6.1	2.3	1.5	6.3	1.3	1.0	5.9		0.0
6.6	3.8	2.5	5.2	2.5	2.0	5.4	2.6	2.5	5.3		1.0
6.6	3.1	2.5	6.2	3.1	1.5	6.4	2.7	1.5			1.0
6.1	0.0	0.0	6.2	0.0	0.0	5.6	2.7	1.5			1.0
5.6	2.8	3.0	5.9	3.7	3.0	6.1	3.9	3.0			1.5
6.9	4.7	3.0	6.1	1.3	0.5	6.4	3.1	•			1.0
6.4	3.0	1.5	6.9	3.8	2.5	6.0	1.5	1.0	6.1	4.0	2.0
	Lesion diam. (cm) (ave.)  5.4  4.8  5.6  6.1  5.6  6.6  6.6  6.7  5.6  6.9  6.4	Lesion diam. (cm) (cm) (ave.)  5.4 0.0  4.8 0.0  5.6 3.6  6.1 3.1  5.6 0.0  6.6 3.8  6.6 3.1  6.1 0.0  5.6 2.8  6.9 4.7	diam. (cm) (cm) (ave.)         tion diam. (cm) density (ave.)           5.4         0.0         0.0           4.8         0.0         0.0           5.6         3.6         2.5           6.1         3.1         1.5           5.6         0.0         0.0           6.6         3.8         2.5           6.1         0.0         0.0           5.6         2.8         3.0           6.9         4.7         3.0           6.4         3.0         1.5	Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Sporulation density (ave.)         Lesion diam. (cm) (ave.)           5.4         0.0         0.0         5.2           4.8         0.0         0.0         4.8           5.6         3.6         2.5         5.3           6.1         3.1         1.5         6.1           5.6         0.0         0.0         6.1           6.6         3.8         2.5         5.2           6.6         3.1         2.5         6.2           6.1         0.0         0.0         6.2           5.6         2.8         3.0         5.9           6.9         4.7         3.0         6.1           6.4         3.0         1.5         6.9	Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (cm) (ave.)         Sporulation diam. (cm) (ave.)           5.4         0.0         0.0         5.2         2.1           4.8         0.0         0.0         4.8         2.8           5.6         3.6         2.5         5.3         2.3           6.1         3.1         1.5         6.1         4.0           5.6         0.0         0.0         6.1         2.3           6.6         3.8         2.5         5.2         2.5           6.6         3.1         2.5         6.2         3.1           6.1         0.0         0.0         6.2         0.0           5.6         2.8         3.0         5.9         3.7           6.9         4.7         3.0         6.1         1.3           6.4         3.0         1.5         6.9         3.8	Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Sporulation diam. (cm) (cm) (ave.)         Sporulation diam. (cm) (cm) (ave.)         Sporulation diam. (cm) (cm) (ave.)           5.4         0.0         0.0         5.2         2.1         1.0           4.8         0.0         0.0         4.8         2.8         2.5           5.6         3.6         2.5         5.3         2.3         1.0           6.1         3.1         1.5         6.1         4.0         2.5           5.6         0.0         0.0         6.1         2.3         1.5           6.6         3.8         2.5         5.2         2.5         2.0           6.6         3.1         2.5         6.2         3.1         1.5           6.1         0.0         0.0         6.2         0.0         0.0           5.6         2.8         3.0         5.9         3.7         3.0           6.9         4.7         3.0         6.1         1.3         0.5	P. capsici OP97         P. capsici SP98         F           Lesion diam. (cm) (ave.) (ave.) (ave.)         Sporulation diam. (cm) (ave.) (ave.)         Lesion diam. (cm) diam. (cm) diam. (cm) (ave.)         Sporulation diam. (cm) diam. (cm) (ave.)         Lesion diam. (cm) diam. (cm) diam. (cm) (ave.)         Lesion diam. diam. (cm) diam. (cm) diam. (cm) (ave.)         Lesion diam. diam. (cm) diam. diam. (cm) diam. (cm) diam. diam. (cm) diam. diam. diam. diam. diam. (cm) diam. diam. (cm) d	P. capsici OP97         P. capsici SP98         P. capsici 23           Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Sporulation diam. (cm) (ave.)         Lesion diam. (cm) (ave.)         Sporulation diam. (cm) (cm) (ave.)         Sporulation diam. (cm) (ave.)         Sporulation diam. (cm) (cm) (cm) (ave.)         Sporulation diam. (cm) (cm) (cm) (cm) (cm) (cm) (cm) (cm)	P. capsici OP97	P. capsici OP97	Lesion diam. (cm) (cm) (ave.)   Component (ave.)

Variety (two fruits per replicate due to limited fruit set on plants, unless	Р.	<i>capsici</i> OP	97	P	. capsici SP	98	F	. capsici 23	8		P. capsici 2	36
indicated otherwise)  P=pickling variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Discover, Seminis <sup>P</sup>	5.4	0.0	0.0	5.4	0.0	0.0	5.7	0.0	0.0	6.0	0.0	0.0
PI 163213, USDA	5.0	0.0	0.0	5.7	2.8	2.0	5.8	3.0	1.0	6.1	1.8	\ \
PI 163214, USDA	5.7	3.1	2.5	5.7	3.0	2.5	5.2	0.0	0.0	5.7	2.3	1.0
PI 249562, USDA**	5.0	2.0	1.0	5.4	0.0	0.0	5.5	0.0	0.0	5.8	3.0	3.0
PI 271327, USDA	5.9	0.0	0.0	5.9	3.1	1.5	5.5	0.0	0.0	5.8	0.0	0.0
PI 279466, USDA	6.0	3.8	2.5	6.1	3.1	1.5	6.3	0.0	0.0	5.9	3.3	2.0
PI 279467, USDA	6.7	3.1	2.0	6.3	1.8	0.5	6.1	3.1	2.5	6.5	3.1	1.5
PI 279468, USDA	5.9	0.0	0.0	6.1	1.5	1.0	5.9	1.6	1.0	6.1	0.0	0.0
PI 321009, USDA	5.9	3.3	2.5	6.0	3.3	3.0	5.9	3.8	2.0	5.7	3.0	2.0
PI 390240, USDA	6.3	3.3	2.5	5.6	0.0	0.0	5.6	3.1	1.5	6.3	1.8	0.5
PI 432867, USDA	6.1	3.0	2.0	5.9	2.3	1.5	5.9	3.0	2.0	6.4	3.4	2.0

'Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation. "Only one fruit per replicate.

## Characterization and Epidemiology of *Phytophthora capsici* Populations and Screening for Genetic Resistance to Fruit Rot in Pickles, 2000-2001

#### M. Hausbeck, K. Lamour, and R. Hammerschmidt

Objective 1: Determine the environmental conditions (temperature and relative humidity) required for sporulation of *Phytophthora capsici* on cucumbers.

Temperature study: Four isolates of *Phytophthora capsici* (OP97, 236, 238, and SP98) were grown on unclarified V8 juice agar plates (10 each) in growth chambers maintaining the following four temperatures: 15°, 22°, 27°, and 33°C. These temperatures span the previously reported range of growth temperatures for *P. capsici*. Growth was recorded daily. This experiment was conducted twice and the results averaged together. The average rate of growth differed depending on the isolate, but in all cases optimal growth was achieved at 27°C (Figure 1). Trials at 4° and 8°C resulted in no-growth of any of the isolates.

Relative Humidity (RH) study: Isolate OP97 (A1 compatibility type) was obtained from a naturally infected pickling cucumber fruit in the northwest region of Michigan during 1997. Single zoospore isolation, compatibility type determination, and long term storage were as previously described. Isolate OP97 was inoculated onto, and re-isolated from, a cucumber fruit 2 weeks prior to the initiation of the experiments and maintained on unclarified V8 juice agar (160 ml V8 juice, 3 g CaCO<sub>3</sub>, 16 g agar, and 840 ml distilled water).

Eight slicing type cucumbers (approximately 15 cm long x 5.0 cm in diameter) obtained from a local supermarket were gently washed and immersed in 0.25% sodium hypochlorite for 5 to 10 minutes, rinsed in distilled water and air dried. A 7 mm V8 agar plug containing actively growing *P. capsici* mycelium and an agar plug without mycelium were placed on opposite ends of the intact surface of each fruit. To prevent plugs from drying out, a 1.5 ml microcentrifuge tube without the cap and with the lip of the cap coated with petroleum jelly was placed over the plug for 24 hours.

Experiments were conducted in a growth chamber (Controlled Environments Inc., Pembina, N. D.) that provided 14 hours of light from two 60W cool white fluorescent bulbs and maintained 23.5°C night/25.5°C day temperatures. Inoculated cucumbers were incubated for five days at RH levels of 60%, 80%, and 98%. Relative humidity was maintained using a RH control unit (Cole-Parmer Instrument Co., Vernon Hills, IL), and temperature and RH were recorded every ten minutes with a data logger (HOBO, Spectrum Technologies Inc., Plainfield, IL).

Cucumbers were observed daily via a covered window built into the side of the chamber and the presence of visible symptoms recorded. On the fifth day (approximately 120 hrs) cucumbers were removed from the chamber, lesion diameter measured, and converted to square centimeters. Sporangia were gently dislodged from lesions using a medium toothbrush into 200 ml of a 300 ppm rose bengal solution and counted with a hemacytometer. The estimated number of sporangia/ml was multiplied by 200 and divided by the total square centimeters of the lesion to obtain an estimate of total sporangia production. Each experiment was conducted twice. A three way analysis of variance (SigmaStat) with a balanced design was conducted to detect interactions between sporangia/cm² and trial, RH, and fruit. Small sections were removed from the edge of the lesions at the conclusion of each experiment and examined with a scanning electron microscope as described below.

Phytophthora capsici isolate OP97 developed visible water soaked lesions on cucumber fruit by the third day post-inoculation and visible sporulation by the fourth day at all three RH levels in the chamber experiments. Results from replicate experiments were combined for analysis because of the low level of variation between experimental results. Average lesion diameters at day 5 were 9.9, 10.0 and 10.2 cm for the 98%, 80%, and 60% RH treatments. Analysis of variance indicates a significant interaction between RH and number of sporangia/cm². Pairwise multiple comparison (Student-Newman-Keuls method) indicates that sporangia production at 98% RH was significantly less than at 60% or 80%, whereas sporangia production at 60% and 80% RH were not significantly different (Figure 2). Our results indicate that sporangia production at 60% and 80% RH is significantly greater than at 98% RH. These results are markedly different than the optimal conditions described for most *Phytophthora* species investigated and suggests that ambient RH levels as low as 60% are not be a limiting factor in the production of sporangia by *P. capsici* on cucumber fruit.

Continued characterization of P. capsici's life history is included in Appendix 1.

#### Objective 2: Screen pickling cucumber germplasm for resistance to P. capsici fruit rot.

Cucumbers of 46 different varieties were grown according to standard practices in fields with a negative history for *P. capsici* in 2001. Mature fruit were harvested and stored in a cold room (4°C) until used. Fruit were soaked 5 minutes in a 5% commercial bleach solution and gently washed, rinsed in distilled water, and dried under ambient conditions. A 0.7 mm plug of actively growing OP97 mycelium was placed at the center of labeled fruit. A plastic microcentrifuge tube was placed over the plug and sealed to the fruit with petroleum jelly to maintain high humidity during initial infection of the fruit. Fruit were incubated for 3 to 4 days on a bench top at room temperature and scored for lesion and sporulation diameter, and density of sporulation. Each experiment was conducted twice. See Appendix 2 for tables. Statistical analyses have not yet been conducted to determine whether there are significant differences among the tested varieties.

### APPENDIX 2

### **Objective 2:**

Screen pickling cucumber germplasm for resistance to *P. capsici* fruit rot.

Cultivar (four fruits per replicate),				P. c	capsici O	P97			
incubated 4 days at room temperature	Lesi	on diam.	(cm)	Sporul	ation diar	n. (cm)	Sport	ılation de	nsity*
1	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave
SS-58137, Sun Seeds 2001	5.6	5.6	5.6	3.3	3.4	3.3	2.5	1.8	2.1
SS-58139, Sun Seeds 2001	5.4	5.6	5.5	3.1	2.6	. 2.9	2.8	2.0	2.4
SS-58141, Sun Seeds 2001	5.6	5.8	5.7	2.3	2.4	2.3	1.5	2.3	1.9
SS-58142, Sun Seeds 2001	5.8	6.0	5.9	2.9	2.2	2.6	1.8	2.3	2.0
SS-58143, Sun Seeds 2001	6.0	6.1	6.0	3.0	2.0	2.5	1.8	2.3	2.0
SS-58144, Sun Seeds 2001	5.1	5.4	5.2	2.4	2.0	2.2	1.8	2.0	1.9
SS-58145, Sun Seeds 2001	5.9	6.1	6.0	3.1	3.3	3.2	1.5	1.8	1.6
SS-58146, Sun Seeds 2001	5.0	5.1	5.0	1.5	1.9	1.7	0.8	0.3	0.5
SS-58147, Sun Seeds 2001	5.6	5.7	5.6	2.3	2.4	2.3	1.8	1.0	1.4
SS-58148, Sun Seeds 2001	5.6	5.3	5.4	2.9	2.0	2.4	2.8	2.0	2.4
SS-58149, Sun Seeds 2001	5.3	5.7	5.5	2.0	1.6	1.8	1.8	0.3	1.0
SS-58150, Sun Seeds 2001	4.8	5.1	5.0	2.0	2.0	2.0	1.8	0.5	1.1
SS-58151, Sun Seeds 2001	5.1	5.2	5.1	1.2	0.6	0.9	0.8	1.5	1.1
SS-58152, Sun Seeds 2001	5.2	5.2	5.2	2.1	2.1	2.1	1.8	1.8	1.8
SS-58153, Sun Seeds 2001	5.4	5.4	5.4	2.4	2.4	2.4	2.3	2.0	2.1
SS-58154, Sun Seeds 2001	5.4	5.4	5.4	2.8	2.8	2.8	1.8	2.0	1.9
SS-58155, Sun Seeds 2001	6.5	6.3	6.4	3.1	3.0	3.1	2.3	2.0	2.1
SS-58456, Sun Seeds 2001	5.1	5.0	5.0	3.2	2.5	2.9	3.0	2.8	2.9
Excel M, Asgrow 2000	6.2	6.5	6.4	3.1	3.2	3.1	3.0	2.0	2.5
Arabian, Asgrow 2000	5.3	5.5	5.4	2.1	2.6	2.3	1.8	1.5	1.6
Vlaspik + M, 2000	5.1	5.5	5.3	2.2	2.2	2.2	1.8	2.0	1.9
Stallion, 2000	5.0	5.2	5.1	2.4	1.6	2.0	2.5	1.8	2.1
Discover M, Asgrow 2000	6.3	6.4	6.3	2.5	3.1	2.8	2.3	2.8	2.5

\*Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

Cultivar (four fruits per replicate),				Р. с	capsici O	P97			
incubated 4 days at room - temperature	Lesion diam. (cm)			Sporulation diam. (cm)			Sporulation density*		
	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave
Discover M, Asgrow 2000	7.0	7.1	7.0	3.2	2.0	2.6	2.5	1.8	2.1
PI 249561, USDA 2001	6.3	6.3	6.3	2.8	2.5	2.7	2.8	2.8	2.8
PI 321006, USDA 2001	5.5	5.2	5.3	2.3	2.1	2.2	2.3	2.5	2.4
PI 321007, USDA 2001	5.6	5.5	5.5	3.4	3.3	3.4	2.8	2.8	2.8
PI 321008, USDA 2001	5.1	5.3	5.2	2.9	2.7	2.8	2.0	2.3	2.1
PI 390261, USDA 2001	5.3	5.0	5.2	2.1	1.4	1.7	2.5	1.5	2.0
PI 390262, USDA 2001	6.1	5.8	6.0	3.0	2.3	2.6	3.0	3.0	3.0
PI 401732, USDA 2001	5.8	5.7	5.7	2.8	2:6-	2.7	2.5	2.5	2.5
PI 401733, USDA 2001	6.0	6.0	6.0	3.1	2.8	3.0	2.5	2.5	2.5
WI 5551, 1994	5.1	5.5	5.3	3.0	2.9	3.0	2.5	3.0	2.8

<sup>\*</sup>Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

Cultivar (four fruits per replicate),				Р. с	capsici Ol	P97			* ;	
incubated 3 days* at room temperature	Lesion diam. (cm)			Sporul	Sporulation diam. (cm)			Sporulation density**		
	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	
Discover M, Asgrow 2000	4.0	3.6	3.8	1.6	0.3	0.9	1.0	0.3	0.6	
PI 197085, USDA 2001	2.6	2.4	2.5	1.7	1.8	1.7	0.5	1.0	0.8	
PI 197088, USDA 2001	3.0	2.6	2.8	1.3	1.2	1.3	0.3	1.0	0.6	
PI 249562, USDA 2001	3.2	3.5	3.4	1.1	1.2	1.2	1.0	0.5	0.8	
PI 271326, USDA 2001	3.3	3.2	3.3	1.2	1.4	1.3	0.5	0.8	0.6	
PI 271327, USDA 2001	3.2	3.4	3.3	1.3	1.6	1.5	1.8	1.8	1.8	
WI 1983 G, 1997	3.3	3.3	3.3	1.9	1.4	1.6	2.0	1.3	1.6	
WI 6632 E, 1997	3.2	3.6	3.4	1.7	1.4	1.5	1.3	1.0	1.1	
WI 5207, 2000	3.5	3.6	3.5	1.5	1.7	1.6	1.5	0.8	1.1	
(WI) SMR 18, 2000	3.7	3.5	3.6	1.7	1.3	1.5	1.3	1.0	1.1	
(WI) GY 14, 1998	3.3	3.9	3.6	1.6	1.2	1.4	1.0	0.5	0.8	

\*Experiment incubated three days due to contamination.

<sup>\*\*</sup>Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

For	EPA	Use	Onl
			ID:

#### Worksheet 3-A(15)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Resistant Cultivars	Study: Evaluation of fungicides and host resistance for
	control of Phytophthora crown rot of summer
	squash, 1999.
Section I Initial Screening on T	echnical Fessibility of Alternatives

1. Are there any location-specific restrictions that inhibit the	use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
M		

For EPA Use Only	
ID#	

### Worksheet 3-A(15)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

1. Is the study on EPA's website?	? Yes No X
1a. If not on the EPA we	ebsite, please attach a copy.
2. Author(s) or researcher(s)	G.J. Holmes
	M.E. Lancaster
	F.J. Louws
8. Publication and Date of Publica	Fungicide and Nematicide Tests, 2000
. Location of research study	Hendersonville, North Carolina
. Name of alternative(s) in study. Resistant Cultivars	. If more than one alternative, list the ones you wish to discuss.
-	
Was crop yield measured in the	e study?  Yes NoX  ne alternative in controlling pests in the study.
Was crop yield measured in the	e study? Yes NoX
. Was crop yield measured in the	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by
. Was crop yield measured in the  Describe the effectiveness of the Using a resistant cultivar was not Phytophthora capsici with this con	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by
. Was crop yield measured in the  Describe the effectiveness of the Using a resistant cultivar was not Phytophthora capsici with this con	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by introl measure.
Was crop yield measured in the Describe the effectiveness of the Using a resistant cultivar was not Phytophthora capsici with this con Discuss how the results of the other factors that would affect y	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by introl measure.
Was crop yield measured in the Describe the effectiveness of the Using a resistant cultivar was not Phytophthora capsici with this con Discuss how the results of the other factors that would affect y	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by introl measure.  study apply to your situation. Would you expect similar results? Are there your adoption of this tool?
. Was crop yield measured in the  Describe the effectiveness of the Using a resistant cultivar was not  Phytophthora capsici with this could be compared to the country of	ne alternative in controlling pests in the study.  commercially acceptable, because there was nearly 40% of the plants killed by introl measure.  study apply to your situation. Would you expect similar results? Are there your adoption of this tool?

OMB Control # 2060-0482

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EVALUATION OF FUNGICIDES AND HOST RESISTANCE FOR CONTROL OF PHYTOPHTHORA CROWN ROT OF SUMMER SQUASH, 1999: The experiment was conducted in a commercial squash field near Hendersonville, NC (GPS coordinates: N35°19.078', W082°25.178') where a severe outbreak of Phytophthora crown rot (PCR) was observed on squash in the summer of 1997. The field was planted to sweet corn in 1997. Soil type was a Codorus loam. The PCR-resistant squash variety was a gray zucchini (SSXP210; Harris Moran Seeds). Treatments were randomized in four complete blocks. Plots were two rows on 4-ft centers, 20 ft long and separated by two rows of pepper. Preplant incorporated (PPI) treatments were applied immediately prior to planting using a CO<sub>2</sub> backpack sprayer equipped with a single nozzle, handheld boom, hollow cone nozzle tip (TXVK-8) and operating at 40 psi. Squash was direct seeded on 7 Jun into ridged beds. Foliar treatments were applied using the same apparatus, by a single pass on each side of the plant bed for a total volume of 56 gal/A. Ridomil Gold + Dithane foliar applications were made on 23 Jun, 7 and 20 Jul and 4 Aug. All other treatments except those that were preplant incorporated (PPI) were applied weekly beginning 16 Jun and ending 11 Aug for a total of 9 applications.

Disease incidence (% mortality) evaluations began 7 days following the first observation of disease (approximately 4 wilted plants in entire test site) when plants had approximately 6 true leaves. Disease progressed rapidly with 46% of individual plots showing greater than 80% plant death on 14 Jul. Yield was not evaluated since the main effect of the disease was plant death. Highly significant block and treatment effects were detected (P=0.01) at each evaluation and for Area Under Disease Progress Curve (AUDPC). The Ridomil Gold EC treatment provided superior control of the disease compared with all other chemical treatments. Because disease attacked early, we believe that most of the effect was due to the PPI treatment rather than subsequent foliar applications. The resistant variety also held up relatively well under the intense disease pressure. However, this variety does not possess good marketable characteristics. A postharvest evaluation (10 fruit from each treatment stored for 20 days at room temperature) yielded 2 out of 120 fruit rotting due to P. capsici. Louws et al. report results of the parallel study on pepper in this volume.

				Mortal	ity (%) 1				
Product and amount/A	29 Jun	07 Jul	14 Jul	20 Jul	28 Jul	03 Aug	11 Aug	17 Aug	AUDPC
Acrobat 50WP, 0.4 lb + Dithane 75DF, 2 lb	11.7 abc	47.7 b	56.2 b	73.0 ab	71.8 b	78.5 b	81.1 b	83.0 b	3278 b
Acrobat 50WP, 0.4 lb + Kocide 2000, 1.9 lb	16.9 bc	56.7 b	66.0 b	79.3 abc	86.7 ab	91.8 abc	93.8 bc	94.5 ab	3851 bc
Acrobat 50WP, 0.4 lb + Dithane 75DF, 2 lb (stem base alt.) $^2$	15.6 bc	52.3 b	59.8 b	71.5 b	75.8 b	80.9 bc	82.5 b	84.1 b	3441 bc
Dithane 75DF, 2 lb	22.2 c	58.6 b	64.2 b	81.2 abc	84.1 bc	89.3 abc	88.9 bc	89.7 bc	3862 bc
Ridomil Gold 4EC, 2 pt (PPI) <sup>3</sup> , Ridomil Gold 4EC, 2 pt + Dithane 75DF, 2 lb (foliar)	0.7 a	1,9 a	3.3 a	. 5.6 a	9.8 a	18.4 a	25.6 a	29.4 a	551 a
4 Quadris 2.08F, 2 pt (PPI)	9.1 abc	61.3 b	73.1 b	86.7 abc	91.1 bc	97.1 bc	97.8 bc	98.5 bc	3974 bc
<sup>4</sup> Flint 50WG, 1 lb (PPI)	13.3 abc	68.6 b	74.3 b	91.8 bc	95.9 c	99.2 c	100 с	100 с	4204 bc
<sup>4</sup> Sovran 50WG, 1 lb (PPI)	18.3 bc	68.5 b	80.2 Ъ	88.6 abc	91.7 bc	95.4 abc	96.6 bc	97.0 bc	4207 bc
<sup>4</sup> Acrobat 50WP, 4 lb (PPI)	12.1 с	72.0 ъ	81.2 ъ	86.6 abc	89.3 bc	93.7 abc	94.9 bc	94.9 bc	4235 bc
Resistant variety, no fungicide	0.0 a	6.9 a	11.2 a	22.1 a	23.7 a	28.5 a	30.6 a	36.5 a	1204 a
Resistant variety + Acrobat 50WP, 0.4 lb + Dithane 75DF, 2 lb	5.3 ab	7.6 a	18.7 a	21.0 a	26.9 a	27.5 a	31.8 a	34.6 a	1112 a
Non-treated	19.9 с	61.2 b	74.8 b	94.9 с	98.3 c	100 c	100 c	100 c	4295 c
LSD (P=0.05)	14.1	25.7	26.6	20.0	20.0	17.8	17.5	15.8	980

Values are the means of 4 replicate plots. Treatments followed by the same letter within a column are not significantly different (K=100, Duncan-Waller K-ratio test).

alternated between spray directed at the base of the plant and directed at the entire plant (not possible until plant height was >1.5 ft; approximately 20 Jun).

<sup>&</sup>lt;sup>3</sup> PPI = preplant incorporated in top 4 inches of soil.

<sup>&</sup>lt;sup>4</sup> Followed by foliar treatment at 14 and 28 days after planting.

Incubated 5 days

Variety	P. capsi	ci OP97	P. capsi	ici SP98	P. capsi	ici SFF3	P. caps	rici SF3
	Lesion diam. (cm) (ave) <sup>1</sup>	Sporulation density² (ave)	Lesion diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation density (ave)
ACX 18, Abbott & Cobb	8.2	1.8	8.4	1.0	8.7	1.5	8.2	0.5
ACX 5001, Abbott & Cobb	8.6	1.6	8.9	1.4	8.8	1.4	8.3	0.6
ACX 5002, Abbott & Cobb	8.6	0.6	8.7	1.8	8.6	2.0	8.5	0.3
Dasher II, Petoseed	8.4	0.9	8.6	1.5	8.8	1.4	8.3	0.1
General Lee, Harris Moran Seed	8.5	1.0	8.2	1.3	8.5	1.4	8.3	0.0
Greensleeves, Harris Moran Seed	8.2	2.1	8.3³	$0.8^{3}$	8.83	1.33	8.6³	1.03
Panther, Sun Seeds	9.0	1.3	8.73	1.8	8.6	1.9	8.34	1.44
Speedway, Petoseed	8.5	1.8	8.4	2.6	8.9	1.8	8.2	0.8
SRQS 2387, Sun Seeds	8.2	0.6	8.5	1.6	8.8	1.1	8.8	0.8
SRQS 2389, Sun Seeds	8.9	1.9	8.6 <sup>4</sup>	1.54	9.0	1.3	8.1	0.1
Ultra Pak, Stokes	8.8	0.8	8.83	2.13	8.6	1.4	8.5	0.1
Vlaspick (pickling)	8.4	2.9	8.5	2.9	8.7	2.9	8.3³	$2.5^{3}$

Average of two replications (four fruits per replicate).

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Three fruits averaged in replicate 1.

Two fruits averaged in replicate 1.

### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH**

Experiments 1 and 2, incubated 3 days

Variety	<i>P</i>	. capsici OP9	7		P. capsici SP9	8	/	P. capsici SFF:	3	P. capsici SF3		
	Lesion diam. (cm) (ave) <sup>1</sup>	Sporulation diam. (cm) (ave)	Sporulation density <sup>2</sup> (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm) (ave)	Sporulation density (ave)	Lesion diam. (cm) (ave)	Sporulation diam. (cm) (ave)	Sporulation density (ave)
Dividend, Rogers	4.6	2.8	1.0	4.3	2.5	1.3	4.4	2.8	1.0	4.5	3.5	1.5
Fortune, Rogers	4.6	4.1	1.8	4.9	4.5	2.0	4.5	4.1	1.5	4.4	3.7	2.0
Golden Dawn II, Rogers	4.9	3.7	1.5	4.3	3.0	1.5	4.2	3.0	1.0	4.5	3.6	2.3
Multipak, Harris Moran Seed	4.3	4.1	1.8	4.0	3.4	2.0	4.3	3.8	2.0	4.3	4.0	2.3
Revenue, Rogers	4.7	3.6	1.8	3.9	3.2	1.8	4.9	3.7	2.0	4.5	3.5	1.5
RSQ 496-VP, Rogers	5.2	4.1	2.0	5.0	3.8	1.8	5.2	4.0	2.5	5.8	4.7	2.3
RSQ 7703-VP, Rogers	4.5 <sup>4</sup>	3.74	2.34	5.2	3.3	1.3	4.6	3.3	1.8	4.94	3.84	2.0⁴
RSQ 8057, Rogers	4.8	3.9	1.5	5.0	3.8	2.0	4.7	3.4	1.5	5.5	3.9	2.0
RSQ 8058, Rogers	4.3	3.1	1.5	4.8	3.7	1.8	5.6	4.1	1.8	$4.5^{3}$	$3.5^{3}$	1.5³
RSQ 8067, Rogers	5.4	4.3	2.0	5.1	4.1	2.3	5.6	4.2	2.0	5.1	3.5	1.8
Spineless Beauty, Rogers	4.7	3.5	2.0	4.9	3.0	1.5	5.5	3.9	1.8	5.1	4.0	1.8
Tigress, Harris Moran Seed	5.5	4.4	2.3	4.5	4.0	2.0	5.1	3.6	1.3	5.2	3.8	1.8
Zucchini Elite, Harris Moran Seed	4.5	3.6	2.5	4.8	3.7	1.8	4.9	3.6	2.0	5.0	4.0	2.3

Average of two replications (two fruits per replicate).

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

One fruit in replicate 1.

One fruit in replicate 2.

## 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici SQUASH Experiment 3, incubated 4 days

Variety (two fruits per replicate)		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Cougar, Harris Moran Seed	4.0	3.4	2.5	4.5	3.7	2.0	4.1	3.0	1.5	4.3	3.4	2.0
General Patton, Asgrow	4.2	3.5	2.0	4.4	4.0	2.5	4.6	3.3	2.0	3.9	3.4	2.5
Golden Rod, Harris Moran Seed	4.5	3.7	2.0	4.7	0.0	0.0	4.2	0.0	0.0	4.2	0.0	0.0
HMX 8714, Harris Moran Seed	4.7	2.6	1.5	4.0	1.6	1.0	4.1	0.0	0.0	4.7	2.8	2.0
HMX 8714, Harris Moran Seed	3.3	2.2	2.0	4.3	1.6	1.5	3.9	0.0	0.0	4.4	1.4	1.0
HMX 8727, Harris Moran Seed	3.5	1.3	0.5	4.0	2.7	1.0	4.1	1.3	0.5	3.4	1.0	0.5
HMX 9705, Harris Moran Seed	2.8	0.7	0.5	3.0	0.0	0.0	2.3²	0.0 <sup>2</sup>	$0.0^{2}$	4.0	1.3	0.5
HMX 9706, Harris Moran Seed	3.1	1.2	1.0	3.5	2.9	1.5	2.3	1.0	0.5	3.2	1.3	1.0
Liberator II, Asgrow	5.3	3.9	1.5	4.4	3.0	1.5	4.3	2.5	1.5	4.9	3.6	2.0
Medallion, Abbott & Cobb	4.0	3.5	1.5	4.6	3.7	1.5	4.4	2.8	1.0	4.2	3.2	2.0
Revenue, Rogers	3.3	0.0	0.0	3.0	1.6	1.0	3.1	0.0	0.0	1.3	0.0	0.0
SSXP 709, Harris Moran Seed	3.7	2.8	2.0	3.6	2.7	1.5	3.7	3.0	2.5	3.8	3.8	2.0
SSXP 787, Harris Moran Seed	5.2	3.1	1.5	5.0	3.0	2.0	3.2	0.0	0.0	4.3	1.8	1.5
SSXP 788, Harris Moran Seed	4.7	2.5	1.5	4.9	3.0	1.5	4.1	0.0	0.0	4.7	1.9	1.0

## 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici SQUASH Experiment 3, incubated 4 days continued

SSXP 789, Harris Moran Seed	4.1	3.1	1.5	4.8	2.8	2.5	3.5	0.6	0.5	3.0	2.0	1.5
SSXP 798, Harris Moran Seed	3.5	2.4	1.0	3.6	2.3	1.0	3.6	1.6	1.0	3.5	2.3	1.5
Superpik, Harris Moran Seed	3.7	3.4	3.0	4.4	3.1	3.0	4.2	3.2	1.5	4.1	3.1	2.5
Superset, Harris Moran Seed	4.0	3.1	1.5	3.8²	2.82	1.0²	4.6	3.0	1.0	3.5	2.7	2.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. )ne fruit per replicate.

# 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to *Phytophthora capsici* SQUASH

Experiment 4, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (avc.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Cougar, Harris Moran Seed	4.9	4.1	1.5	5.0	3.2	2.0	4.9	3.3	1.5	4.2	2.3	1.5
General Patton, Asgrow	4.9	4.4	2.5	5.0	5.0	2.0	4.6	4.6	1.0	5.5	5.2	1.5
Golden Rod, Harris Moran Seed	3.6	3.4	1.0	3.72	2.72	$2.0^{2}$	4.6	3.4	1.0	3	3	<sup>3</sup>
HMX 8714, Harris Moran Seed	5.2	3.5	2.5	4.7	3.4	2.0	5.4	1.5	1.0	5.0	3.4	2.0
HMX 8714, Harris Moran Seed	3.62	2.5 <sup>2</sup>	2.0 <sup>2</sup>	5.4	3.4	2.0	5.6	3.1	2.0	5.4 <sup>2</sup>	3.72	$3.0^{2}$
HMX 8727, Harris Moran Seed	4.3	4.3	1.0	4.7²	$3.9^{2}$	2.0 <sup>2</sup>	4.5 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$	4.6²	2.82	1.0 <sup>2</sup>
HMX 9705, Harris Moran Seed	4.2²	4.22	1.02	3.3²	3.3 <sup>2</sup>	2.0 <sup>2</sup>	3	3	3	3	3	3
HMX 9706, Harris Moran Seed	4.2	3.8	1.5	4.7	3.8	1.5	4.6	3.4	1.5	4.7	4.0	1.5
Liberator II, Asgrow	5.3	4.9	2.0	5.7	5.5	1.5	5.8	5.3	2.0	5.5	5.5	2.0
Medallion, Abbott & Cobb	4.2	4.2	2.0	5.0	4.6	2.0	5.7	5.2	3.0	4.7	4.0	3.0
Revenue, Rogers	4.1	3.1	1.0	5.7	4.8	1.0	5.1	3.7	1.5	5.9	4.6	2.0
SSXP 709, Harris Moran Seed	4.5	4.1	3.0	4.8	4.5	2.0	6.0	5.3	2.5	5.7	5.4	2.5
SSXP 787, Harris Moran Seed	5.3	3.1	1.5	5.4	2.3	1.0	5.2	3.2	1.5	5.3	4.1	2.0
SSXP 788, Harris Moran Seed	5.0	2.9	1.0	3.9	3.1	2.5	5.0	2.9	1.5	4.7	4.7	1.5

### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH** Experiment 4, incubated 3 days

#### continued

SSXP 789, Harris Moran Seed	4.9	3.6	3.0	5.4	3.7	2.0	4.5	3.4	2.5	5.0	2.7	1.5
SSXP 798, Harris Moran Seed	3.9	3.1	2.0	4.6	4.1	1.0	4.9	3.1	2.0	4.1	3.0	1.0
Superpik, Harris Moran Seed.	4.7	3.6	2.0	4.9	3.2	1.5	4.5	3.5	1.0	4.8	4.6	2.0
Superset, Harris Moran Seed .	5.7	5.4	2.5	5.5	5.5	2.5	5.7	3.4	1.5	5.7	4.4	1.5

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

<sup>&</sup>lt;sup>2</sup>One fruit per replicate.

<sup>3</sup>Fruits were too contaminated to evaluate.

## 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici SQUASH Experiment 5, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
ACX 34, Abbott & Cobb	5.7	4.6	2.5	6.12	3.5 <sup>2</sup>	2.02	5.4	4.2	2.0	5.7	4.4	3.0
Dividend VIP, Siegers Seed	5.1	4.2	3.0	5.9	5.0	2.0	6.4	4.6	2.0	5.7	4.5	3.0
Golden Dawn	4.6	3.6	1.5	4.5	3.8	1.5	4.9	3.4	1.0	4.5	3.4	2.0
Revenue, Siegers Seed	5.9	4.7	2.0	5.7	4.4	2.0	5.5	4.1	2.0	5.3	4.5	2.5
3XS 9732, Sun Seeds	5.8	4.5	3.0	5.5	3.8	2.0	5.6	3.8	2.0	5.3	4.3	2.5
XRHT 1777	4.2	3.6	2.0	4.1	3.0	1.0	4.6	4.1	2.0	4.8	3.8	2.5
Zucchini Elite F1, Harris Moran Seed	5.2	3.9	2.0	5.5	4.5	2.0	4.7	3.2	2.0	4.7	3.5 .	. 2.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. )ne fruit per replicate.

# 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to *Phytophthora capsici* SQUASH

Pickling cucumber Experiment 4, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3	P. capsici SF3		
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
ACX 34, Abbott & Cobb	4.9	3,5	2.0	4.8	3.3	2.5	4.5²	3.0 <sup>2</sup>	2.0 <sup>2</sup>	5.0	3.3	2.0
Dividend VIP, Siegers Seed	3.6	0.6	0.5	5.0	3.9	2.0	4.0	1.5	0.5	4.6	3.0	2.0
Revenue, Siegers Seed	4.5	3.2	2.0	4.8	3.4	2.0	5.2	3.7	2.0	4.6	2.8	1.5
Seasons, Abbott & Cobb	3.6	1.9	1.5	5.2	1.8	0.5	5.9	3.6	2.0	5.4	3.8	1.5
Seneca Prolific, Siegers Seed .	4.9	4.1	2.0	5.4	4.5	1.5	5.3	4.1	2.0	4.6	1.8	0.5
SXS 9732, Sun Seeds	5.6	4.2	3.0	5.4	3.9	1.5	5.0	3.7	2.0	4.9	3.2	2.5
Zucchini Elite F1, Harris Moran Seed	5.2	3.3	1.0	4.2	3.3	1.5	5.3	3.6	2.0	4.0	1.8	1.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

One fruit per replicate.

### 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH**Rerun of pickling cucumber Experiment 4, incubated 3 days

Variety (two fruits per		P. capsici OP9	7	P. capsici SP98				P. capsici SFF	3	P. capsici SF3			
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density ' (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	
ACX 34, Abbott & Cobb	4.6	3.8	1.5	5.1	3.8	1.0	4.8	3.8	1.5	2	2	2	
Dividend VIP, Siegers Seed	4.5	3.3	2.5	4.7	4.1	1.0	4.8	3.7	1.5	4.2	1.9	0.5	
Revenue, Siegers Seed	3.8 .	1.7	1.0	4.4	1.7	0.5	4.8	1.7	1.0	3.6	1.4	1.0	
Seasons, Abbott & Cobb	4.0	1.8	1.0	5.1	4.3	1.0	4.5	3.9	1.0	3.5	1.6	0.5	
Seneca Prolific, Siegers Seed .	4.8	3.9	3.0	4.7	4.2	2.5	4.7	3.8	2.5	3.8	3.3	2.0	
SXS 9732, Sun Seeds	4.9	3.7	2.5	4.4	3.3	1.0	2.5	1.3	0.5	4.4	3.4	1.5	
Zucchini Elite F1, Harris Moran Seed	4.6	3.2	1.5	4.2	2.7	1.0	4.0	1.5	0.5	4.5	2.0	1.5	

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Fruits were too contaminated to evaluate.

# 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to *Phytophthora capsici* SQUASH

Pickling cucumber Experiment 6, incubated 3 days

Variety (two fruits per		P. capsici OP9	7	P. capsici SP98				P. capsici SFF	3	P. capsici SF3		
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
13367, USDA	5.4	3.9	1.0	3.8	0.0	0.0	3.9	0.0	0.0	5.6	1.0	0.5

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

# 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to *Phytophthora capsici* SQUASH

Pickling cucumber Experiment 7 (redo's), incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
13367, USDA	3.5	2.6	1.5	4.1	2.2	1.	5.3	3.2	1.0	4.3	2.5	1.0
Goldfinger	5.0 <sup>2</sup>	3.32	1.02	3	3	3	4.2²	3.3 <sup>2</sup>	$1.0^{2}$	3.5	2.3	1.0
Golden Dawn III	3	3	3	3	<b></b> <sup>3</sup> .	3	4.2²	4.2 <sup>2</sup>	1.02	3.1 <sup>2</sup>	2.0 <sup>2</sup>	1.0 <sup>2</sup>
XRHT 1777	5.5	4.9	2.0	4.6	3.7	2.0	- 5.1	4.3	1.5	<sup>'</sup> 4.5	3.3	1.5

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Fruits were too contaminated to evaluate.

One fruit per replicate.

Experiment 1, incubated 6 days

'ariety (four fruits per		P. capsici OP9	07		P. capsici SP9	8		P. capsici SFF	3		P. capsici SI	73
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
lalypso, Atlas Seeds	9.1	8.6	3.0	8.3⁴	5.24	3.04	9.6	8.7	3.0	8.7³	7.2 <sup>3</sup>	$3.0^{3}$
arolina, Atlas Seeds	9.8	8.3	3.0	9.5⁴	7.54	3.04	10.1	8.1	3.0	8.7	7.2	2.8
Cross Country F1, Harris Aoran Seed	9.0	8.6	3.0	8.03	6.13	$3.0^{3}$	9.6	8.8	3.0	8.83	6.23	2.73
Discover M Hybrid, Asgrow .	9.7	8.2	3.0	10.1	6.8	3.0	10.6	7.7	3.0	11.3	7.8	3.0
X 1914 183491, Seminis	9.1	8.6	3.0	7.7²	4.82	2.52	9.5	8.1	3.0	9.4	8.9	3.0
excel M, Asgrow	9.5	7.8	3.0	8.6⁴	5.44	2.04	11.6	8.1	3.0	8.7³	7.0 <sup>3</sup>	$3.0^{3}$
ancipak, Asgrow	10.1	9.1	3.0	8.9	6.3	3.0	10.9	9.1	3.0	10.3	8.4	3.0
MX 5020 F1, Harris Moran eed	9.93	8.43	$3.0^{3}$	7.5²	4.9²	2.5 <sup>2</sup>	9.8	7.6	3.0	8.4	6.1	2.8
IMX 3469 F1, Harris Moran	9.5	7.1	3.0	7.6²	4.0 <sup>2</sup>	1.52	9.8	8.2	3.0	9.03	5.9 <sup>3</sup>	2.33
IMX 8460 F1, Harris Moran	10.4	8.2	3.0	9.3	6.4	2.8	10.7	9.0	3.0	9.7	7.6	3.0
IMX 8461 F1, Harris Moran leed	10.9	8.9	3.0	9.2	6.5	3.0	10.4	9.4	3.0	9.5	8.0	3.0
afayette Classic, Sun Seeds .	10.1	8.4	3.0	7.4	4.8	2.8	9.4	8.6	3.0	10.7	8.0	3.0
'I 209064, USDA	10.1	8.9	3.0	7.6²	4.8 <sup>2</sup>	$3.0^{2}$	9.8	8.3	3.0	9.4	8.7	3.0
<sup>2</sup> I 426169, USDA	10.1	7.3	3.0	8.5⁴	6.24	3.04	10.0	7.6	3.0	10.0	6.6	2.8
'I 466922, USDA	9.6	7.7	3.0	8.8	6.4	3.0	9.6	8.3	3.0	10.3	8.0	3.0
Pioneer, Atlas Seeds	9.4	8.3	3.0	7.6²	6.1 <sup>2</sup>	$3.0^{2}$	9.9	8.8	3.0	9.8	6.5	2.5

Experiment 1, incubated 6 days Continued

/ariety (four fruits per eplicate)		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	73
ерпсассу	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
legal F1, Harris Moran Seed.	10.4	8.4	3.0	9.2³	7.4 <sup>3</sup>	$3.0^{3}$	10.8	8.6	3.0	10.6	8.4	3.0
loyal Fl, Harris Moran Seed.	9.6	8.8	3.0	8.9³	$7.5^{3}$	$3.0^{3}$	9.3	8.6	3.0	10.2	8.4	3.0
3RQP 2391, Sun Seeds	10.4	9.0	3.0	5	5	<b></b> 5	9.2	8.5	3.0	8.7	7.8	3.0
Stallion 193782, Seminis Seed	9.9	8.5	3.0	10.7	8.0	3.0	10.3	9.5	3.0	11.4	9.5	3.0
Sumter, Atlas Seeds	9.9	7.8	3.0	9.12	6.2 <sup>2</sup>	$3.0^{2}$	11.0	8.0	3.0	9.7	7.2	3.0
Famor Hybrid, Asgrow	10.7	9.5	3.0	10.4	8.7	3.0	10.4	9.2	3.0	10.0	9.1	3.0
Fransamerica F1, Harris Moran Geed	9.7	7.7	3.0	8.5²	$7.0^{2}$	3.02	10.8	8.4	3.0	9.2	8.8	3.0
Victoria, Sun Seeds	10.3	8.1	3.0	9.4	9.0	3.0	10.3³	9.23	$3.0^{3}$	11.4	9.8	3.0
√laspik VGA733, Seminis	10.2	7.5	3.0	9.3	5.7	2.5	10.7	7.4	3.0	9.1	6.5	2.5
√lasset, Asgrow	8.7	8.1	3.0	8.7	6.7	3.0	9.9	8.7	3.0	9.2	8.2	3.0
√lasspear Hybrid, Asgrow	9.6	8.4	3.0	8.8³	$6.6^{3}$	$3.0^{3}$	10.1	9.0	3.0	9.1	8.2	3.0
√lasstar B, Asgrow	9.2	7.6	3.0	6.9²	3.9 <sup>2</sup>	1.5 <sup>2</sup>	10.4	8.4	3.0	9.1	7.0	3.0
Wisconsin, Atlas Seeds	9.6	7.4	3.0	9.0²	8.2 <sup>2</sup>	3.0 <sup>2</sup>	10.3	8.2	3.0	10.9	8.4	3.0

porulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

wo fruits per replicate.

hree fruits per replicate.

ne fruit per replicate.

.ll fruits were too contaminated to evaluate.

Experiment 2, incubated 3 days

Variety (four fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Carolina, Atlas Seeds	3.9³	2.83	2.0³	4.2	2.6	1.8	4.6	1.6	1.0	3.8	2.3	1.5
Cross Country F1, Harris Moran Seed	4.9³	3.33	2.03	3.5	2.4	1.33	4.8	2.5	1.3	4.3	3.0	1.5
Discover M Hybrid, Asgrow .	4.7	1.6	1.0	4.2	1.8	1.0	5.0 <sup>2</sup>	$3.0^{2}$	1.5 <sup>2</sup>	5.1	1.8	1.3
EX 1911 155633, Seminis	4.5	2.9	2.0	4.1	2.6	2.0	4.3 <sup>2</sup>	3.4 <sup>2</sup>	$2.0^{2}$	4.23	2.83	1.33
EX 1914 183491, Seminis	4.7³	3.4 <sup>3</sup>	$2.0^{3}$	4.5³	$3.3^{3}$	$2.0^{3}$	3.8	3.1	2.0	4.5²	3.3 <sup>2</sup>	2.5 <sup>2</sup>
Excel M, Asgrow	4.2	2.7	1.5	4.8	3.0	1.8	5.0 <sup>3</sup>	$3.1^{3}$	1.73	4.3³	$2.9^{3}$	1.73
HMX 3469 F1, Harris Moran Seed	4.5³	3.43	$2.0^{3}$	3.72	3.12	2.0 <sup>2</sup>	4.44	2.84 .	1.04	6.0²	4.3 <sup>2</sup>	3.02
HMX 8460 F1, Harris Moran Seed	5.2	3.4	2.0	4.9	2.8	1.5	4.6²	3.5 <sup>2</sup>	2.02	4.8	2.8	1.8
HMX 8461 F1, Harris Moran Seed	4.1	3.0	1.8	4.4	2.9	1.73	4.7³	3.13	1.33	· 4.5³	3.13	$2.0^{3}$
Jackson, Sun Seeds	4.9	3.9	2.0	5.2	2.2	1.3	4.9	3.4	2.0	5.3³	4.43	2.73
Lafayette Classic, Sun Seeds .	5.0	4.1	2.3	4.43	$2.8^{3}$	$2.0^{3}$	5.83	$2.7^{3}$	1.73	4.6³	3.33	$2.0^{3}$
PI 466922, USDA	5.0 <sup>2</sup>	3.4 <sup>2</sup>	1.52	4.1 <sup>2</sup>	2.22	$1.5^{2}$	4.8 <sup>2</sup>	$3.6^2$	2.0 <sup>2</sup>	3.9 <sup>3</sup>	$1.9^{3}$	1.33
Regal F1, Harris Moran Seed .	3.9 <sup>3</sup>	$3.4^{3}$	$2.0^{3}$	4.8 <sup>3</sup>	$3.3^{3}$	1.73	5.2 <sup>3</sup>	3.23	$1.3^{3}$	4.2³	$2.7^{3}$	1.73
Royal F, Harris Moran Seed1.	4.13	2.83	$1.0^{3}$	4.5³	3.13	$2.0^{3}$	4.5³	$3.0^{3}$	1.73	3.6 <sup>2</sup>	2.1 <sup>2</sup>	1.52
Stallion 193782, Seminis	4.6 <sup>2</sup>	3.1 <sup>2</sup>	$2.0^{2}$	2.54	2.54	2.04	3.33	$2.8^{3}$	1.73	4.6	3.5	2.0
Tamor Hybrid, Asgrow	4.2	2.5	1.5	4.1 <sup>2</sup>	$2.8^{2}$	$1.5^{2}$	4.7³	3.43	$2.0^{3}$	3.2 <sup>2</sup>	2.4 <sup>2</sup>	1.02
Vlasstar B, Asgrow	4.5	3.1	2.0	3.5	1.8	1.0	4.9	2.2	1.0	4.5	3.0	1.8

Experiment 2, incubated 3 days Continued

Variety (four fruits per replicate)		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
Tophodicy	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density! (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Wisconsin, Atlas Seeds	5.4	2.1	1.0	4.4	3.0	1.8	4.9²	2.9²	1.5 <sup>2</sup>	5.2	3.1	1.8

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. I'wo fruits per replicate.

Three fruits per replicate.

One fruit per replicate.

Experiment 3, incubated 4 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Calypso, Atlas Seeds	6.0	4.1	2.0	6.3	4.5	1.5	6.9	4.9	2.5	5.9	3.7	2.0
EX 1911 155633, Seminis	6.3	4.3	2.0	5.4	4.9	2.0	5.9	4.7	2.0	7.0	4.8	2.0
Fancipak, Asgrow	6.9	3.9	1.5	7.6	4.1	2.0	6.3	5.1	2.0	5.9	3.4	1.5
FMX 5020 F1, Harris Moran Seed	6.6	5.0	2.5	6.8	4.9	2.0	6.6	5.1	2.5	6.1	5.3	2.5
PI 209064, USDA	6.9	4.0	1.5	6.9	5.2	2.0	6.4	4.4	2.0	6.4	4.3	1.5
PI 390241, USDA	6.1 <sup>2</sup>	6.1 <sup>2</sup>	$3.0^{2}$	6.1	5.6	3.0	5.5 <sup>2</sup>	6.3 <sup>2</sup>	1.0 <sup>2</sup>	5.5	4.9	2.5
PI 391570, USDA	5.9²	. 4.6 <sup>2</sup>	$2.0^{2}$	4.5	3.9	2.0	5.7	4.9	2.0	3.3²	3.3 <sup>2</sup>	$2.0^{2}$
PI 422182, USDA	6.2	4.8	2.0	5.8	1.8	0.5	6.8	5.6	2.5	6.8²	3.7 <sup>2</sup>	$2.0^{2}$
PI 426169, USDA	5.9	3.9	1.5	5.5	2.3	1.0	6.1	4.2	1.5	5.9	5.0	2.0
PI 426170, USDA	5.8	3.9	1.5	5.7	3.4	1.5	6.0	4.2	2.0	6.0	3.3	1.0
PI 432890, USDA	5.1	4.6	2.0	5.2	3.3	1.5	5.6²	5.6 <sup>2</sup>	$2.0^{2}$	6.4	5.0	2.0
PI 483339, USDA	5.6 <sup>2</sup>	$3.8^{2}$	$2.0^{2}$	6.0	4.4	1.5	5.3	4.3	2.0	4.2	3.4	2.0
Pioneer, Atlas Seeds	6.5	4.7	4.0 .	5.9	2.5	1.5	7.7	5.2	1.0	4.7	4.7	2.0
SRQS 2389, Sun Seeds	5.2	5.2	2.5	6.4	5.8	2.5	5.6	4.7	2.5	7.1	5.5	2.0
SRQP 2391, Sun Seeds	4.7 <sup>2</sup>	4.72	$2.0^{2}$	4.2²	4.2 <sup>2</sup>	$2.0^{2}$	6.6	5.1	3.0	7.0	4.0	2.0
Transamerica F1, Harris Moran Seed	6.0	4.2	2.0	6.0	4.1	2.0	5.1	3.7	1.5	5.5	4.1	2.0
Vlaspik VGA733, Seminis	6.7	2.3	1.0	5.8	3.6	2.0	7.0	4.9	2.0	6.3	2.9	1.5
Vlasset, Asgrow	5.5	5.5	3.0	5.3	4.9	2.5	5.9	5.0	2.0	6.0	4.1	2.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

One fruit per replicate.

Experiment 4, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>(</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
PI 167223, USDA	3,7	2.8	1.5	4.8	3.0	1.0	4.3	2.5	1.0	3.3 <sup>2</sup>	0.02	0.0²
PI 197086, USDA	3.4	2.1	1.0	3.8 <sup>2</sup>	2.72	$2.0^{2}$	4.4	2.1	1.0	3.2²	1.62	1.0 <sup>2</sup>
PI 209069, USDA	3.5	0.0	0.0	3.4	0.0	0.0	4.9	2.5	1.5	3.7	1.0	0.5
PI 234517, USDA	5.0	0.0	0.0	4.3 <sup>2</sup>	1.92	$2.0^{2}$	3	3	3	1.5²	1.5 <sup>2</sup>	1.02
PI 271328, USDA	5.0 <sup>2</sup>	2.12	1.0 <sup>2</sup>	3	3	3	3	3	3	3.3²	$0.0^{2}$	0.02
PI 288238, USDA	3	3	3	5.0 <sup>2</sup>	3.4 <sup>2</sup>	$2.0^{2}$	5.3 <sup>2</sup>	3.12	2.0 <sup>2</sup>	4.0	2.8	1.5
PI 330628, USDA	4.3	2.1	1.5	4.0	1.2	0.5	4.6	3.0	1.5	4,4	1.5	0.5
PI 390241, USDA	3.4	1.2	0.5	3.4	2.0	1.0	4.3	2.1	1.0	3	3	<sup>3</sup>
PI 391570, USDA	3.6²	$0.0^{2}$	$0.0^{2}$	3	3	<b></b> 3	4.1 <sup>2</sup>	2.12	1.02	4.2²	$3.0^{2}$	1.0 <sup>2</sup>
PI 422182, USDA	4.1	1.9	1.0	4.6	2.3	1.0	4.5	2.7	1.0	4.0	2.2	1.0
PI 426170, USDA	4.0	1.8	1.0	$3.7^{2}$	$0.0^{2}$	$0.0^{2}$	4.2²	$0.0^{2}$	$0.0^{2}$	2.1 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
PI 432851, USDA	3	3	3	3	3	3	3	3	3	3	<b></b> 3	3
PI 432855, USDA	3.6²	2.5	$2.0^{2}$	3	3	3	3	3	3	3	3	3
PI 483339, USDA	4.4	1.2	0.5	4.7²	2.9 <sup>2</sup>	1.02	3	3	3	3.6²	2.22	1.0 <sup>2</sup>
SRQS 2389, Sun Seeds	4.1	2.7	2.0	4.0	2.2	1.5	4.0	2.2	1.5	3.8	2.8	1.5

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. One fruit per replicate.

All fruits were too contaminated to evaluate.

Experiment 5, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
PI 163213, USDA	4.6	2.3	1.0	4.9	2.5	1.0	5.2	2.2	1.0	4.2	2.3	1.0
PI 167223, USDA	5.4	3.1	1.5	5.6	3.5	1.0	8.1	3.5	1.0	5.0	1.5	0.5
PI 197086, USDA	3.1	0.5	0.5	2.6	2.6	1.0	3.4	2.2	1.5	4.0	2.6	2.0
PI 197088, USDA	3.4	1.0	0.5	5.0	3.2	1.5	3.5	0.0	0.0	3.4	0.0	0.0
PI 209069, USDA	5.4	3.3	1.5	4.9	3.0	1.5	4.6	2.6	1.0	4.7	2.2	1.5
PI 227209, USDA	4.6	2.1	1.0	5.2	3.1	1.5	4.8	2.4	1.0	5.3	3.3	1.5
PI 234517, USDA	4.8	2.9	1.5	3.9	1.3	0.5	3.9	0.9	0.5	3.8	1.4	0.5
PI 267942, USDA	4.6	2.9	1.5	5.2	3.2	1.0	4.8	2.1	1.0	4.2	2.2	1.0
PI 271328, USDA	3.7	0.0	0.0	5.3	1.3	0.5	3.7	1.2	0.5	4.1	0.0	0.0
PI 288238, USDA	5.4	3.2	0.5	5.7	3.7	2.0	5.6	1.7	1.5	4.9	2.7	1.0
PI 330628, USDA	4.2	2.7	2.0	4.5	2.5	1.5	4.3	1.3	0.5	2.8	0.0	0.0
PI 390244, USDA	4.6	2.6	1.5	4.9	2.8	1.0	4.7	0.0	0.0	4.4	2.1	1.0
PI 390529, USDA	5.0	1.9	1.0	4.2	1.9	1.0	5.7	3.1	2.0	3.9	2.0	1.0
PI 418964, USDA	5.1	2.8	1.5	5.0	2.7	1.5	4.5	2.2	1.0	3.6	2.7	1.0
PI 432851, USDA	5.8	3.7	2.0	4.6	2.5	1.0	4.2	1.4	1.0	4.6	2.5	0.5
PI 432855, USDA	5.5	3.4	3.0	5.5	3.2	1.5	5.8	3.1	1.5	4.9	2.8	1.5
PI 432865, USDA	5.7	4.5	3.0	4.8	3.7	2.0	4.4	2.6	1.0	4.5	3.0	1.5
PI 432890, USDA	4.5	2.7	1.0	4.7	1.7	1.0	6.3	1.7	1.0	5.0	2.4	1.0
Sumter, Atlas Seeds	4.7	3.4	2.0	5.0	3.2	1.5	5.5	2.4	1.0	4.9	2.8	2.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Experiment 6, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
eplicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density 1 (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Ames 7118, USDA	4.5	3.4	3.0	4.0	3.0	1.0	4.6	1.5	0.5	3.7	1.2	0.5
lackson, Sun Seeds	5.1	3.3	2.5	4.7	3.2	1.0	5.7	3.7	1.5	4.2	2.3	2.0
PI 163213, USDA	4.4	1.1	0.5	5.0	1.3	0.5	4.8	0.0	0.0	4.9	0.0	0.0
PI 197088, USDA	3.5²	$2.0^{2}$	1.02	4.0	1.3	0.5	3.7	0.0	0.0	4.3	0.8	0.5
PI 211979, USDA	5.7	2.9	2.0	5.2	1.4	0.5	4.5	0.0	0.0	4.4	0.7	0.5
PI 249562, USDA	4.1	0.0	0.0	4.3	1.0	0.5	4.9	0.8	0.5	4.6	0.0	0.0
PI 267942, USDA	4.3	0.9	0.5	5.4	2.0	1.0	4.1	1.2	0.5	3.3	0.0	0.0
PI 279466, USDA	5.1	2.2	1.0	5.2	1.2	0.5	4.9	1.9	1.0	3.9²	$0.0^{2}$	$0.0^{2}$
PI 279467, USDA	4.4	0.7	0.5	4.1	0.0	0.0	3.1	0.0	0.0	4.4	1.0	0.5
PI 279468, USDA	4.8	1.0	0.5	4.8	1.1	0.5	3.7	0.0	0.0	2.9	0.0	0.0
PI 321008, USDA	5.9	0.0	0.0	5.5	1.2	0.5	5.2	0.0	0.0	4.8	0.9	0.5
PI 390240, USDA	5.0²	$0.0^{2}$	$0.0^{2}$	4.7²	$0.0^{2}$	$0.0^{2}$	6.3	1.5	0.5	5.2 <sup>2</sup>	1.52	1.0 <sup>2</sup>
PI 390244, USDA	4.4	1.9	1.0	5.2	0.7	0.5	4.5	1.0	0.5	5.0	0.8	0.5
PI 390246, USDA	5.4	2.1	1.0	5.6	2.4	1.0	5.6	2.7	1.0	4.3	0.0	0.0
PI 390262, USDA	3.7	2.4	1.0	3.6	2.0	1.0	5.2	2.3	1.0	4.5²	$2.0^{2}$	1.02
PI 390529, USDA	4.9	2.4	1.0	4.7	1.1	0.5	5.0	0.0	0.0	5.2	1.0	0.5
PI 418964, USDA	5.2	2.2	1.0	6.2	2.8	1.0	4.7	2.4	1.0	3.8	0.8	0.5
PI 432865, USDA	4.6	3.1	3.0	3.8	1.9	1.0	5.8	2.9	1.0	4.8	2.9	2.0
PI 432867, USDA	5.5	2.9	1.0	5.4	2.9	1.0	5.1	1.0	0.5	4.9	0.0	0.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. One fruit per replicate.

Experiment 7 (redo's), incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
Ames 7118, USDA	4.6	3.3	2.0	4.0	2.9	1.5	4.2	2.5	1.5	4.0	3.1	2.0
Excel M, Asgrow	4.2	0.0	0.0	5.6	0.0	0.0	4.5	0.0	0.0	3.6	0.0	0,0
HMX 3469 F1, Harris Moran Seed	4.8	0.6	0.5	4.5	0.0	0.0	4.4	0.0	0.0	3.2	0.0	0.0
PI 211979, USDA	5.2	2.8	1.5	5.3	3.1	1.0	4.7	3.4	1.0	4.6	2.3	1.0
PI 279468, USDA	5.1	4.6	1.5	4.6	3.3	2.5	4.9	3.8	2.0	4.6	3.7	2.0
PI 390241, USDA	5.2	3.8	3.0	5.3	4.0	2.0	4.0	3.4	2.0	4.3	3.1	1.0
PI 390246, USDA	5.0	2.9	1.0	5.1	3.0	1.0	4.7	2.4	1.0	6.1	4.4	1.5
PI 422182, USDA	3.9	0.0	0.0	4.3	0.0	0.0	4.0	0.0	0.0	1.9	0.0	0.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

Rerun of Experiment 4, incubated 3 days

Variety (two fruits per	-	P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>(</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
PI 167223, USDA	4.9	3.4	2.5	4.6	2.6	1.0	4.4	3.2	1.5	4.8	3.3	2.0
PI 197086, USDA	4.0	2.5	2.0	3.8	1.2	0.5	4.0	2.2	1.5	3.2	0.0	0.0
PI 209069, USDA	3.9	0.0	0.0	4.5	1.2	0.5	3.9	1.8	1.0	3.7	0.0	0.0
PI 234517, USDA	3.3	3.3	3.0	4.2	2.9	1.0	3.8	2.9	2.0	2.8	0.0	0.0
PI 271328, USDA	3.6	0.0	0.0	4.2	2.2	1.0	4.1	0.0	0.0	3.6	1.4	0.5
PI 288238, USDA	5.2	3.9	1.5	4.7	3.7	2.0	4.9	3.8	3.0	4.3	3.1	2.5
PI 330628, USDA	4.1	2.4	2.0	4.5	1.2	0.5	4.5	2.8	1.5	3.5	0.0	0.0
PI 391570, USDA	4.6	2.8	1.0	4.5	2.3	1.0	3.3	0.0	0.0	3.9	3.0	2.0
PI 426170, USDA	4.5	1.3	0.5	4.3	3.0	1.5	4.5	0.0	0.0	3.6	1.5	0.5
PI 432851, USDA	4.7	2.8	2.0	4.8	3.6	3.0	4.7	1.4	0.5	4.3	1.6	0.5
PI 432855, USDA	4.0	3.5	2.0	4.4	3.1	2.0	4.7	3.4	2.5	4.2	3.0	1.0
PI 483339, USDA	4.5	2.3	1.5	4.6	3.4	1.5	3.4	1.4	1.0	4.6	1.7	0.5
SRQS 2389, Sun Seeds	3.3	0.9	1.0	5.1 <sup>2</sup>	2.72	1.0²	3.2	1.7	1.0	2.6	0.0	0.0

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy. One fruit per replicate.

Experiment 8, incubated 3 days

Variety (two fruits per		P. capsici OP9	7		P. capsici SP9	8		P. capsici SFF	3		P. capsici SF	3
replicate)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)	Lesion diam. (cm) (average)	Sporulation diam. (cm) (average)	Sporulation density <sup>1</sup> (average)
Calypso, Atlas Seeds	4.5	1.6	0.5	4.4	3.2	2.0	4.4	3.3	1.5	4.8	2.6	1.5
Carolina, Atlas Seeds	3.9	3.5	3.0	4.7	4.0	2.5	4.3	3.8	1.0	4.2	3.7	2.5
PI 249562, USDA	3	3	3	<b></b> 3	3	3	3	3	3	3	3	3
PI 279466, USDA	3	3	3	4.2 <sup>2</sup>	3.22	1.02	3	3	3	4.0²	2.5 <sup>2</sup>	2.0 <sup>2</sup>
PI 279467, USDA	<sup>3</sup>	3	3	3.9²	$3.9^{2}$	$2.0^{2}$	3	3	3	3	3	3
PI 358813, USDA	<b></b> 3	3	3	4.6²	$2.0^{2}$	$1.0^{2}$	3	3	3	3	3	3
PI 390240, USDA	3.9²	3.22	1.02	3	3	3	3.9²	$2.5^{2}$	$2.0^{2}$	3.5²	2.7 <sup>2</sup>	$2.0^{2}$
PI 390241, USDA	5.1	3.8	1.5	4.8	3.6	2.5	4.2	3.1	1.0	5.4²	$4.0^{2}$	2.0 <sup>2</sup>
PI 422182, USDA	4.0	3.0	2.0	4.9	3.8	2.0	6.0	3.9	1.0	4.5	3.4	2.0
PI 432867, USDA	3	3	3	3	3	3	4.1	3.4	1.0	3	3	3
PI 432890, USDA	4.6²	$3.0^{2}$	1.0 <sup>2</sup>	3	3	<b>7-</b> 3	5.0²	2.2 <sup>2</sup>	$1.0^{2}$	4.7	3.1	1.5
PI 483339, USDA	4.5²	$2.5^{2}$	2.0 <sup>2</sup>	5.1 <sup>2</sup>	3.5 <sup>2</sup>	$1.0^{2}$	4.6²	3.8 <sup>2</sup>	1.02	2.5 <sup>2</sup>	$0.0^{2}$	$0.0^{2}$
SRQP 2391, Sun Seeds	4.1	3.7	2.5	4.0	4.0	2.0	4.6	4.1	2.0	$4.0^{2}$	4.0 <sup>2</sup>	2.0 <sup>2</sup>
Stallion 193782, Seminis	3.2 <sup>2</sup>	3.22	1.0²	3.4 <sup>2</sup>	3.4 <sup>2</sup>	2.0 <sup>2</sup>	3	3	3	4.0²	$4.0^{2}$	$2.0^{2}$

Sporulation density visually rated on a score of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy.

One fruit per replicate.

All fruits were too contaminated to evaluate.

## 1999 MICHIGAN STATE UNIVERSITY GERM PLASM TRIAL: Resistance to Phytophthora capsici **SQUASH**Run of Experiment 7 (redo's), incubated 3 days

Variety (two fruits per replicate)	P. capsici OP97			P. capsici SP98			P. capsici SFF3			P. capsici SF3		
replicate)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density <sup>1</sup> (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)	Lesion diam. (cm) (ave.)	Sporulation diam. (cm) (ave.)	Sporulation density (ave.)
72 13367	3.5	2.6	. 1.5	4.1	2.2	1.	5.3	3.2	1.0	4.3	2.5	1.0
Goldfinger	5.0²	3.3 <sup>2</sup>	1.02				4.2².	3.3 <sup>2</sup>	1.0 <sup>2</sup>	3.5	2.3	1.0
XRHT 1777	5.5	4.9	2.0	4.6	3.7	2.0	5.1	4.3	1.5	4.5	3.3	1.5
Golden Dawn III			~=				4.2²	4.2 <sup>2</sup>	1.0 <sup>2</sup>	3.1 <sup>2</sup>	2.0²	1.02

One fruit per replicate.

# Screening Cucurbits for Genetic Resistance to Fruit Rot in Pickles, 1999-2000

Submitted by:
M. Hausbeck
R. Hammerschmidt

Characteristics of the challenge inoculum: Four field isolates of *P. capsici* exhibiting diversity for mating type, sensitivity to mefenoxam, host type, origin and AFLP fingerprint were selected as representative of the diversity in Michigan cucurbit production fields. These include: 1) OP97, isolated from pickling cucumber fruit in northwestern Michigan in 1997, A1 mating type, and fully sensitive to mefenoxam; 2) SP98, isolated from pumpkin fruit in southwestern Michigan in 1998, A2 mating type, and fully sensitive to mefenoxam; 3) 238, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A2 mating type, and intermediately sensitive to mefenoxam; 4) 236, isolated from pickling cucumber fruit in southcentral Michigan in 1998, A1 mating type, and sensitive to mefenoxam; 5) a control was included that consisted of an agar plug only with no pathogen present.

Mating type was determined by mating each isolate to known A1 and A2 isolates on unclarified V8 agar (UCV8) plates and scoring for the presence or absence of oospores after a three to five day incubation period. Mefenoxam sensitivity was determined in vitro by placing a 0.7 mm plug of actively expanding mycelium onto the center of 100 x 15 mm UCV8 plates amended with 0 and 100 ppm mefenoxam. Plates were incubated at 23 to 25°C for three days and colony diameter measured. Percent growth on the amended plates was determined relative to the unamended control. Percent growth <30% of the control is designated as sensitive, between 30 to 90% as intermediate sensitivity, and >90% as fully insensitive.

Growing of fruit: Some cucumber cultivars grown in 1999 were selected on the basis of the last year's screening to be grown again in 2000, along with new varieties. Five seed lots of two cultivars ('Reisenschal' and 'Vlaspik') were selected on the basis of different seed treatments. Cucumbers were grown according to standard practices in fields with a negative history for *P. capsici* infection. Mature fruit were harvested weekly, sorted according to size, and stored in a cold room at 4°C until a chamber experiment could be initiated (generally three to six days).

Fruit preparation and inoculation: Fruit were subjected to a 5 minute immersion in a 5% commercial bleach solution and gently washed, then rinsed in distilled water. Fruit were allowed to dry under ambient conditions. Dry fruit were labeled with a numerical code indicating the inoculum and cucumber variety with a permanent marker. A 0.7 mm plug of actively expanding mycelium or plain UCV8 agar was placed at the center of unwounded fruit. A plastic microcentrifuge tube was placed over the agar plug and sealed to the fruit with petroleum jelly to maintain high humidity during initial infection of the fruit.

Experimental design: A completely randomized design determined the layout of inoculated

fruit. A random number generator was used to construct a linear array of the number set used to code individual fruit. Fruit were incubated for four days on a bench top at room temperature and scored for lesion diameter, sporulation density, and the diameter of sporulation.

Each of the five treatment/host combinations was replicated four times in Experiments 1 through 5. Due to a lack of fruit set of some cultivars, there were only two replications per experiment in the last two runs (Experiments 6,7). Each treatment/host combination was represented in at least two experiments.

Number of cucumber cultivars screened for resistance to *Phytophthora capsici* in 2000.

Cucumber type	Number screened
Pickling	21
Slicing	4
Plant introduction	32
Total	57

Further analysis on this year's data will be conducted to choose lines that will be evaluated in next year's screen. While all lines appeared to be susceptible, we are interested in pursuing those lines where lesion diameter and sporulation density was reduced. See Appendix II for tables.

Variety (four fruits per replicate)	Р.	capsici OP	97	Р.	capsici SP	98	F	P. capsici 23	8	I	P. capsici 23	36
P=pickling variety S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Arabian Hybrid, Asgrow <sup>P</sup>	6.7	0.5	0.3	6.7	0.5	0.3	6.9	0.6	0.8	6.6	0.5	0.3
Colt Hybrid, Asgrow <sup>P</sup>	6.7	0.0	0.0	7.0	0.6	0.5	6.8	0.0	0.0	6.9	1.1	0.8
Cyclone, Asgrow <sup>s</sup>	7.6	1.9	1.5	6.7	0.0	0.0	7.5	3.8	2.5	7.2	0.0	0.0
Discover, Seminis <sup>P</sup>	6.9	0.0	0.0	6.8	0.0	0.0	6.9	0.0	0.0	6.7	0.0	0.0
Eureka, Siegers <sup>P</sup>	6.3	1.9	1.3	6.0	0.0	0.0	6.7	2.4	1.8	6.6	1.4	1.0
Excel, Seminis <sup>P</sup>	7.1	1.8	1.3	7.0	0.6	0.5	7.0	0.0	0.0	6.7	0.0	0.0
Lafayette, Sun Seeds <sup>P</sup>	6.8	1.1	0.8	6.5	1.5	1.0	7.0	1.3	1.0	6.9	1.1	1.0
Lightning, Asgrow <sup>s</sup>	7.1	2.4	1.0	7.2	1.4	0.8	7.4	0.0	0.0	7.4	0.6	0.5
Palomino Hybrid, Asgrow <sup>P</sup>	7.3	1.6	1.3	7.4	0.5	0.3	7.0	2.0	1.5	6.4	1.1	0.8
PI 209067, USDA	6.7	3.4	1.3	6.5	2.3	1.0	6.6	1.6	1.0	6.5	1.6	0.8
PI 249561, USDA	7.1	0.0	0.0	6.7	0.0	0.0	6.5	0.0	0.0	6.8	0.0	0.0
PI 426169, USDA	7.4	1.9	1.5	6.8	0.5	0.3	7.0	2.3	1.3	6.6	1.5	0.8
Reisenschal (control), Vlasic Foods <sup>P</sup>	7.2	3.0	2.3	6.9	3.0	2.3	7.1	0.0	0.0	6.8	0.0	0.0
Stallion Hybrid, Asgrow <sup>P</sup>	6.7	0.6	0.5	7.0	0.5	0.5	7.4	0.0	0.0	7.0	0.6	0.5
Thunder, Asgrow <sup>s</sup>	7.4	1.3	0.8	7.3	0.5	0.5	7.5	0.6	0.5	7.4	0.8	0.5
Transamerica, Sun Seeds <sup>P</sup>	7.0	0.7	0.5	6.6	0.5	0.3	7.1	1.3	0.8	6.7	1.1	0.8
Vlaspik (B-1,SMP), Vlasic Foods <sup>P</sup>	6.1	2.5	2.3	6.7	0.0	0.0	7.4	0.0	0.0	6.5	1.6	1.3
Vlaspik (B-2,SMP,+), Vlasic Foods <sup>P</sup>	6.8	0.0	0.0	6.5	0.0	0.0	7.1	0.0	0.0	6.8	0.0	0.0

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	Р.	capsici SP	98	ŀ	P. capsici 23	8	P. capsici 236		
s=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Vlaspik (B-3,SMP,E), Vlasic Foods <sup>P</sup>	6.7	0.0	0.0	6.5	0.0	0.0	6.9	0.0	0.0	6.6	0.0	0.0
Vlaspik (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	7.1	0.0	0.0	6.8	0.0	0.0	6.8	0.0	0.0	6.3	0.0	0.0
Vlaspik (control), Asgrow <sup>P</sup>	7.2	0.5	0.3	6.7	0.0	0.0	7.3	0.5	0.3	6.6	0.0	0.0
Vlaspik+M Hybrid, Asgrow <sup>P</sup>	6.6	0.8	0.8	7.2	0.0	0.0	7.0	0.0	0.0	7.3	0.6	0.5
Vlasspear, Seminis <sup>P</sup>	6.6	2.5	2.0	7.2	2.0	1.5	7.5	0.0	0.0	6.9	0.0	0.0

Rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	P	capsici SP	98	1	o. capsici 23	8	Į.	P. capsici 23	6
S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Arabian Hybrid, Asgrow <sup>P</sup>	6.8	0.5	0.3	6.6	0.5	0.3	6.9	0.0	0.0	6.9	0.0	0.0
Colt Hybrid, Asgrow <sup>P</sup>	6.7	0.0	0.0	7.0	0.6	0.5	7.0	0.0	0.0	7.3	0.6	0.5
Cyclone, Asgrow <sup>s</sup>	7.2	1.6	1.3	7.1	1.5	1.3	7.1	0.0	-0.0	7.1	3.7	2.3
Discover, Seminis <sup>P</sup>	5.0	0.0	0.0	6.7	0.0	0.0	7.0	0.0	0.0	6.8	0.0	0.0
Eureka, Siegers <sup>p</sup>	7.1	1.8	1.5	6.5	0.5	0.5	6.9	2.5	2.3	6.9	2.4	2.3
Excel, Seminis <sup>P</sup>	6.9	2.4	2.3	7.0	1.8	1.5	7.2	0.9	0.8	6.8	0.0	0.0
Lafayette, Sun Seeds <sup>P</sup>	7.0	1.6	1.0	7.1	0.8	0.5	7.2	1.8	1.5	6.7	0.0	0.0
Lightning, Asgrow <sup>s</sup>	7.3	1.6	1.0	7.0	0.9	0.5	7.0	1.4	1.3	7.4	1.8	1.5
Palomino Hybrid, Asgrow P	7.1	0.8	0.8	6.9	0.0	0.0	6.8	1.3	0.8	6.7	0.9	0.8
PI 249561, USDA	7.1	0.6	0.5	7.2	0.9	0.8	7.1	0.0	0.0	7.0	0.0	0.0
PI 426169, USDA	7.1	1.4	1.3	6.8	0.8	0.5	7.1	1.5	1.0	6.9	1.8	1.5
Reisenschal (control), Vlasic Foods <sup>P</sup>	7.3	3.8	2.8	7.1	1.6	1.5	7.1	3.7	3.0	7.2	2.0	1.5
Stallion Hybrid, Asgrow <sup>P</sup>	7.1	0.8	0.5	6.7	0.0	0.0	7.0	0.0	0.0	7.2	0.8	0.8
Thunder, Asgrow <sup>s</sup>	7.3	1.0	0.8	7.1	1.8	1.0	7.4	1.0	0.5	7.3	0.0	0.0
Transamerica, Sun Seeds <sup>P</sup>	7.1	1.5	1.0	6.7	0.6	0.5	6.9	1.5	1.5	6.8	0.5	0.5
Vlaspik (B-1,SMP), Vlasic Foods <sup>P</sup>	6.8	2.5	2.3	6.4	0.8	0.8	6.3	2.5	2.3	6.7	1.0	0.8
Vlaspik (B-2,SMP,+), Vlasic Foods <sup>P</sup>	6.9	0.0	0.0	7.3	0.0	0.0	6.8	0.0	0.0	7.1	0.0	0.0
Vlaspik (B-3,SMP,E), Vlasic Foods <sup>P</sup>	6.7	0.0	0.0	6.6	0.0	0.0	6.7	0.0	0.0	7.1	0.0	0.0

Variety (four fruits per replicate)	P. capsici OP97			P. capsici SP98			I	P. capsici 23	8	P. capsici 236		
=pickling variety =slicing variety  Vlaspik (B-4,SMP,E,+), Vlasic Foods	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Vlaspik (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	6.9	0.0	0.0	6.8	0.0	0.0	6.7	0.0	0.0	7.0	0.0	0.0
Vlaspik (control), Vlasic Foods <sup>P</sup>	7.0	1.6	1.3	7.1	0.9	0.8	7.1	2.7	2.3	7.0	1.7	1.5
Vlaspik+M Hybrid, Asgrow <sup>P</sup>	7.0	0.0	0.0	6.8	1.5	1.3	7.0	0.9	0.8	6.6	0.0	0.0
Vlasspear, Seminis <sup>P</sup>	6.9	1.3	0.8	6.9	0.0	0.0	7.2	0.0	0.0	6.7	0.0	0.0

<sup>\*</sup>Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	Р.	capsici SP	98	I	. capsici 23	38	F	P. capsici 23	36
picking variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Discover, Seminis <sup>P</sup>	6.6	0.0	0.0	6.6	0.0	0.0	6.4	0.0	0.0	6.6	0.6	0.3
PI 209067, USDA	6.2	3.0	2.0	5.6	2.4	1.5	6.0	3.0	2.3	5.6	2.4	1.8
PI 209068, USDA	6.4	- 3.3	1.7	6.5	2.4	1.5	6.3	2.5	1.5	6.1	1.3	0.8
PI 209069, USDA	5.8	3.3	2.3	6.2	3.0	2.5	6.1	3.6	2.5	6.2	2.7	2.0
PI 211980, USDA	5.9	3.8	2.8	6.2	3.9	2.5	6.1	3.4	2.8	6.0	4.0	2.5
PI 271328, USDA	5.3	2.5	1.8	5.2	2.0	1.3	6.0	2.3	2.0	5.6	2.5	2.3
PI 330628, USDA	5.6	2.3	1.5	5.2	2.3	1.5	5.3	3.4	2.3	5.6	3.3	2.5
PI 358813, USDA	6.2	2.4	1.5	6.0	2.7	1.5	6.1	2.8	1.8	6.0	2.1	1.0
PI 358814, USDA	5.9	2.0	1.3	5.1	2.0	1.3	6.4	1.3	0.8	4.0	0.0	0.0
PI 390262, USDA	6.2	3.8	2.5	6.7	4.1	2.8	6.4	3.6	2.5	6.0	3.3	2.3
PI 390263, USDA	5.8	4.0	2.5	6.0	3.8	2.8	5.6	3.7	3.0	6.3	4.1	3.0
PI 426170, USDA	5.6	1.4	1.0	5.0	1.9	1.3	6.0	1.9	1.3	5.4	2.3	1.3
Reisenschal (B-1,SMP), Vlasic Foods <sup>P</sup>	6.1	2.9	1.8	6.6	3.6	2.5	6.4	3.2	2.0	6.6	2.8	2.3
Reisenschal (B-2,SMP,+), Vlasic Foods <sup>P</sup> .	6.5	3.3	2.3	6.2	2.4	1.5	6.4	2.3	1.3	5.6	2.9	2.0
Reisenschal (B-3,SMP,E), Vlasic Foods <sup>P</sup> .	6.4	2.8	2.0	6.1	2.6	1.8	6.3	3.0	2.3	6.4	3.1	2.8
Reisenschal (B-4,SMP,E,+), Vlasic Foods <sup>P</sup>	5.9	5.0	1.8	5.3	2.5	2.0	5.7	2.1	1.5	5.8	2.0	1.3

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	Р.	capsici SP	98	I	P. capsici 23	18	F	. capsici 23	36
s=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Discover, Seminis <sup>P</sup>	5.9	0.0	0.0	6.3	0.0	0.0	6.0	0.0	0.0	5.7	0.0	0.0
Meteor, Asgrow <sup>s</sup>	7.3	0.0	0.0	7.5	0.5	0.3	8.0	1.1	0.8	7.6	0.0	0.0
PI 197087, USDA	5.3	1.0	0.5	6.1	1.4	0.8	6.0	3.9	2.5	6.4	3.6	2.3
PI 197088, USDA	5.7	1.8	1.3	6.0	0.6	0.5	6.1	0.5	0.5	5.4	1.8	1.3
PI 209068, USDA	5.7	0.6	0.5	5.4	1.4	0.8	5.8	0.8	0.5	6.2	1.8	1.0
PI 209069, USDA	6.3	1.5	0.8	6.2	1.3	0.8	6.2	1.8	1.5	6.2	0.6	0.5
PI 211978, USDA	6.8	1.6	1.0	6.4	1.6	1.0	7.0	2.9	1.3	7.0	3.6	2.0
PI 211979, USDA	6.4	3.7	2.3	6.8	3.1	2.3	7.5	3.8	1.8	7.0	2.6	1.5
PI 211980, USDA	6.1	2.0	1.3	6.2	1.4	1.0	6.2	1.1	0.8	6.1	0.8	0.5
PI 271328, USDA	6.2	1.1	0.8	6.2	1.4	1.0	6.1	2.4	1.8	5.9	0.5	0.3
PI 279468, USDA	7.6	3.9	2.8	7.3	2.9	2.0	7.1	2.1	2.0	7.4	3.0	2.3
PI 330628, USDA	6.0	1.4	1.0	6.2	2.4	1.8	5.8	0.0	0.0	6.7	2.8	1.8
PI 358813, USDA	5.9	0.6	0.5	6.6	2.0	1.5	6.8	1.5	1.0	6.6	1.6	1.0
PI 358814, USDA	6.1	1.4	1.0	6.2	0.8	0.3	6.5	1.5	1.0	6.7	2.5	2.0
PI 390262, USDA	6.4	1.3	0.8	6.3	0.8	0.5	6.0	1.4	1.3	6.6	0.8	0.8
PI 390263, USDA	6.7	0.6	0.5	6.1	2.5	1.8	6.0	0.0	0.0	6.1	0.6	0.5
PI 426170, USDA	5.9	0.6	0.5	6.0	1.1	0.8	5.6	2.2	1.3	6.0	1.3	1.0
PI 432868, USDA	7.2	3.3	2.5	7.1	2.8	2.0	7.2	3.1	2.0	6.7	2.8	1.8

Variety (four fruits per replicate)  P=pickling variety	P. capsici OP97			P. capsici SP98			ŀ	P. capsici 23	38	P. capsici 236		
S=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Reisenschal (B-1,SMP), Vlasic Foods <sup>P</sup>	5.7	0.5	0.3	6.1	0.0	0.0	5.7	0.6	0.3	6.4	0.0	0.0
Reisenschal (B-2,SMP,+), Vlasic Foods <sup>P</sup> .	5.9	0.0	0.0	6.2	0.0	0.0	6.0	0.0	0.0	6.1	0.0	0.0
Reisenschal (B-3,SMP,E), Vlasic Foods <sup>P</sup> .	5.9	0.0	0.0	5.6	0.0	0.0	6.0	0.0	0.0	6.1	0.4	0.3
Reisenschal (B-4,SMP,E,+), Vlasic Foods <sup>P</sup> *Snorulation density rated on a scale of 0 to 3 a	5.6	0.0	0.0	6.2	0.0	0.0	6.2	0.0	0.0	6.3	0.0	0.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (four fruits per replicate)  P=pickling variety	Р.	capsici OP	97	P	. capsici SP	98	I	P. capsici 23	8	P. capsici 236		
s=slicing variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Discover, Seminis <sup>P</sup>	6.1	0.0	0.0	5.9	0.0	0.0	6.0	0.0	0.0	6.4	0.0	0.0
Meteor, Asgrow <sup>s</sup>	6.9	0.9	0.3	7.0	1.5	0.8	7.0	2.9	1.3	6.4	0.0	0.0
PI 197088, USDA	6.5	0.6	0.3	6.8	1.4	0.8	5.8	1.8	1.0	6.3	0.8	0.5
PI 21 1978, USDA	6.3	0.6	0.5	6.8	3.1	1.8	6.5	2.3	1.8	6.7	0.6	0.3
PI 227210, USDA	6.8	8.0	0.3	6.9	2.1	1.8	6.8	2.3	1.3	7.0	0.8	0.3
PI 279468, USDA	6.6	1.6	0.8	6.2	2.2	1.3	5.8	0.8	0.5	5.8	3.1	1.8
PI 390239, USDA	6.8	1.5	1.3	6.3	2.2	1.0	5.8	1.6	0.5	6.3	2.3	1.8
PI 432868, USDA	6.2	1.6	1.0	5.6	2.4	1.5	6.3	0.0	0.0	6.4	1.5	1.0

'Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (two fruits per replicate due to	Р.	capsici OP	97	P	. capsici SP	98	ŀ	. capsici 23	8	P. capsici 236		
limited fruit set on plants) <sup>P</sup> ≕pickling variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)
Discover, Seminis <sup>P</sup>	5.4	0.0	0.0	5.2	2.1	1.0	5.1	0.0	0.0	4.9	0.0	0.0
PI 163213, USDA	4.8	0.0	0.0	4.8	2.8	2.5	4.1	0.0	0.0	4.7	0.0	0.0
PI 163214, USDA	5.6	3.6	2.5	5.3	2.3	1.0	5.5	3.0	2.0	4.9	2.3	2.0
PI 227209, USDA	6.1	3.1	1.5	6.1	4.0	2.5	6.1	3.1	1.5	5.9	0.0	0.0
PI 271327, USDA	5.6	0.0	0.0	6.1	2.3	1.5	6.3	1.3	1.0	5.9	0.0	0.0
PI 279466, USDA	6.6	3.8	2.5	5.2	2.5	2.0	5.4	2.6	2.5	5.3	3.0	1.0
PI 279467, USDA	6.6	3.1	2.5	6.2	3.1	1.5	6.4	2.7	1.5	6.1	3.1	1.0
PI 321008, USDA	6.1	0.0	0.0	6.2	0.0	0.0	5.6	2.7	1.5	5.9	2.5	1.0
PI 321009, USDA	5.6	2.8	3.0	5.9	3.7	3.0	6.1	3.9	3.0	6.1	2.3	1.5
PI 390240, USDA	6.9	4.7	3.0	6.1	1.3	0.5	6.4	3.1	2.0	5.7	2.2	1.0
PI 432867, USDA	6.4	3.0	1.5	6.9	3.8	2.5	6.0	1.5	1.0	6.1	4.0	2.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

Variety (two fruits per replicate due to limited fruit set on plants, unless	Р.	capsici OP	97	P	. capsici SP	98	1	P. capsici 23	8	P. capsici 236		
indicated otherwise)  P=pickling variety	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density' (ave.)	Lesion diam. (cm) (ave.)	Sporula- tion diam. (cm) (ave.)	Sporula- tion density* (ave.)
Discover, Seminis <sup>P</sup>	5.4	0.0	0.0	5.4	0.0	0.0	5.7	0.0	0.0	6.0	0.0	0.0
PI 163213, USDA	5.0	0.0	0.0	5.7	2.8	2.0	5.8	3.0	1.0	6.1	1.8	0.5
PI 163214, USDA	5.7	3.1	2.5	5.7	3.0	2.5	5.2	0.0	0.0	5.7	2.3	1.0
PI 249562, USDA"	5.0	2.0	1.0	5.4	0.0	0.0	5.5	0.0	0.0	5.8	3.0	3.0
PI 271327, USDA	5.9	0.0	0.0	5.9	3.1	1.5	5.5	0.0	0.0	5.8	0.0	0.0
PI 279466, USDA	6.0	3.8	2.5	6.1	3.1	1.5	6.3	0.0	0.0	5.9	3.3	2.0
PI 279467, USDA	6.7	3.1	2.0	6.3	1.8	0.5	6.1	3.1	2.5	6.5	3.1	1.5
PI 279468, USDA	5.9	0.0	0.0	6.1	1.5	1.0	5.9	1.6	1.0	6.1	0.0	0.0
PI 321009, USDA	5.9	3.3	2.5	6.0	3.3	3.0	5.9	3.8	2.0	5.7	3.0	2.0
PI 390240, USDA	6.3	3.3	2.5	5.6	0.0	0.0	5.6	3.1	1.5	6.3	1.8	0.5
PI 432867, USDA	6.1	3.0	2.0	5.9	2.3	1.5	5.9	3.0	2.0	6.4	3.4	2.0

<sup>&#</sup>x27;Sporulation density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, and 3=heavy sporulation.

"Only one fruit per replicate.

# Characterization and Epidemiology of *Phytophthora capsici* Populations and Screening for Genetic Resistance to Fruit Rot in Pickles, 2000-2001

#### M. Hausbeck, K. Lamour, and R. Hammerschmidt

Objective 1: Determine the environmental conditions (temperature and relative humidity) required for sporulation of *Phytophthora capsici* on cucumbers.

Temperature study: Four isolates of *Phytophthora capsici* (OP97, 236, 238, and SP98) were grown on unclarified V8 juice agar plates (10 each) in growth chambers maintaining the following four temperatures: 15°, 22°, 27°, and 33°C. These temperatures span the previously reported range of growth temperatures for *P. capsici*. Growth was recorded daily. This experiment was conducted twice and the results averaged together. The average rate of growth differed depending on the isolate, but in all cases optimal growth was achieved at 27°C (Figure 1). Trials at 4° and 8°C resulted in no growth of any of the isolates.

Relative Humidity (RH) study: Isolate OP97 (A1 compatibility type) was obtained from a naturally infected pickling cucumber fruit in the northwest region of Michigan during 1997. Single zoospore isolation, compatibility type determination, and long term storage were as previously described. Isolate OP97 was inoculated onto, and re-isolated from, a cucumber fruit 2 weeks prior to the initiation of the experiments and maintained on unclarified V8 juice agar (160 ml V8 juice, 3 g CaCO<sub>3</sub>, 16 g agar, and 840 ml distilled water).

Eight slicing type cucumbers (approximately 15 cm long x 5.0 cm in diameter) obtained from a local supermarket were gently washed and immersed in 0.25% sodium hypochlorite for 5 to 10 minutes, rinsed in distilled water and air dried. A 7 mm V8 agar plug containing actively growing *P. capsici* mycelium and an agar plug without mycelium were placed on opposite ends of the intact surface of each fruit. To prevent plugs from drying out, a 1.5 ml microcentrifuge tube without the cap and with the lip of the cap coated with petroleum jelly was placed over the plug for 24 hours.

Experiments were conducted in a growth chamber (Controlled Environments Inc., Pembina, N. D.) that provided 14 hours of light from two 60W cool white fluorescent bulbs and maintained 23.5°C night/25.5°C day temperatures. Inoculated cucumbers were incubated for five days at RH levels of 60%, 80%, and 98%. Relative humidity was maintained using a RH control unit (Cole-Parmer Instrument Co., Vernon Hills, IL), and temperature and RH were recorded every ten minutes with a data logger (HOBO, Spectrum Technologies Inc., Plainfield, IL).

Cucumbers were observed daily via a covered window built into the side of the chamber and the presence of visible symptoms recorded. On the fifth day (approximately 120 hrs) cucumbers were removed from the chamber, lesion diameter measured, and converted to square centimeters. Sporangia were gently dislodged from lesions using a medium toothbrush into 200 ml of a 300 ppm rose bengal solution and counted with a hemacytometer. The estimated number of sporangia/ml was multiplied by 200 and divided by the total square centimeters of the lesion to obtain an estimate of total sporangia production. Each experiment was conducted twice. A three way analysis of variance (SigmaStat) with a balanced design was conducted to detect interactions between sporangia/cm² and trial, RH, and fruit. Small sections were removed from the edge of the lesions at the conclusion of each experiment and examined with a scanning electron microscope as described below.

Phytophthora capsici isolate OP97 developed visible water soaked lesions on cucumber fruit by the third day post-inoculation and visible sporulation by the fourth day at all three RH levels in the chamber experiments. Results from replicate experiments were combined for analysis because of the low level of variation between experimental results. Average lesion diameters at day 5 were 9.9, 10.0 and 10.2 cm for the 98%, 80%, and 60% RH treatments. Analysis of variance indicates a significant interaction between RH and number of sporangia/cm². Pairwise multiple comparison (Student-Newman-Keuls method) indicates that sporangia production at 98% RH was significantly less than at 60% or 80%, whereas sporangia production at 60% and 80% RH were not significantly different (Figure 2). Our results indicate that sporangia production at 60% and 80% RH is significantly greater than at 98% RH. These results are markedly different than the optimal conditions described for most *Phytophthora* species investigated and suggests that ambient RH levels as low as 60% are not be a limiting factor in the production of sporangia by *P. capsici* on cucumber fruit.

Continued characterization of P. capsici's life history is included in Appendix 1.

#### Objective 2: Screen pickling cucumber germplasm for resistance to P. capsici fruit rot.

Cucumbers of 46 different varieties were grown according to standard practices in fields with a negative history for *P. capsici* in 2001. Mature fruit were harvested and stored in a cold room (4°C) until used. Fruit were soaked 5 minutes in a 5% commercial bleach solution and gently washed, rinsed in distilled water, and dried under ambient conditions. A 0.7 mm plug of actively growing OP97 mycelium was placed at the center of labeled fruit. A plastic microcentrifuge tube was placed over the plug and sealed to the fruit with petroleum jelly to maintain high humidity during initial infection of the fruit. Fruit were incubated for 3 to 4 days on a bench top at room temperature and scored for lesion and sporulation diameter, and density of sporulation. Each experiment was conducted twice. See Appendix 2 for tables. Statistical analyses have not yet been conducted to determine whether there are significant differences among the tested varieties.

# **APPENDIX 2**

# **Objective 2:**

Screen pickling cucumber germplasm for resistance to *P. capsici* fruit rot.

Cultivar (four fruits per replicate),				Р. с	apsici O	P97			
incubated 4 days at room temperature	Lesi	on diam.	(cm)	Sporul	ation dian	n. (cm)	Sport	ılation de	nsity*
	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave
SS-58137, Sun Seeds 2001	5.6	5.6	5.6	3.3	3.4	3.3	2.5	1.8	2.1
SS-58139, Sun Seeds 2001	5.4	5.6	5.5	3.1	2.6	2.9	2.8	2.0	2.4
SS-58141, Sun Seeds 2001	5.6	5.8	5.7	2.3	2.4	2.3	1.5	2.3	1.9
SS-58142, Sun Seeds 2001	5.8	6.0	5.9	2.9	2.2	2.6	1.8	2.3	2.0
SS-58143, Sun Seeds 2001	6.0	6.1	6.0	3.0	2.0	2.5	1.8	2.3	2.0
SS-58144, Sun Seeds 2001	5.1	5.4	5.2	2.4	2.0	2.2	1.8	2.0	1.9
SS-58145, Sun Seeds 2001	5.9	6.1	6.0	3.1	3.3	3.2	1.5	1.8	1.6
SS-58146, Sun Seeds 2001	5.0	5.1	5.0	1.5	1.9	1.7	0.8	0.3	0.5
SS-58147, Sun Seeds 2001	5.6	5.7	5.6	2.3	2.4	2.3	1.8	1.0	1.4
SS-58148, Sun Seeds 2001	5.6	5.3	5.4	2.9	2.0	2.4	2.8	2.0	2.4
SS-58149, Sun Seeds 2001	5.3	5.7	5.5	2.0	1.6	1.8	1.8	0.3	1.0
SS-58150, Sun Seeds 2001	4.8	5.1	5.0	2.0	2.0	2.0	1.8	0.5	1.1
SS-58151, Sun Seeds 2001	5.1	5.2	5.1	1.2	0.6	0.9	0.8	1.5	1.1
SS-58152, Sun Seeds 2001	5.2	5.2	5.2	2.1	2.1	2.1	1.8	1.8	1.8
SS-58153, Sun Seeds 2001	5.4	5.4	5.4	2.4	2.4	2.4	2.3	2.0	2.1
SS-58154, Sun Seeds 2001	5.4	5.4	5.4	2.8	2.8	2.8	1.8	2.0	1.9
SS-58155, Sun Seeds 2001	6.5	6.3	6.4	3.1	3.0	3.1	2.3	2.0	2.1
SS-58456, Sun Seeds 2001	5.1	5.0	5.0	3.2	2.5	2.9	3.0	2.8	2.9
Excel M, Asgrow 2000	6.2	6.5	6.4	3.1	3.2	3.1	3.0	2.0	2.5
Arabian, Asgrow 2000	5.3	5.5	5.4	2.1	2.6	2.3	1.8	1.5	1.6
Vlaspik + M, 2000	5.1	5.5	5.3	2.2	2.2	2.2	1.8	2.0	1.9
Stallion, 2000	5.0	5.2	5.1	2.4	1.6	2.0	2.5	1.8	2.1
Discover M, Asgrow 2000	6.3	6.4	6.3	2.5	3.1	2.8	2.3	2.8	2.5

\*Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

Cultivar (four fruits per replicate),	P. capsici OP97								
incubated 4 days at room temperature	Lesi	Lesion diam. (cm)		Sporul	ation diar	n. (cm)	Sporulation density*		
•	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave
Discover M, Asgrow 2000	7.0	7.1	7.0	3.2	2.0	2.6	2.5	1.8	2.1
PI 249561, USDA 2001	6.3	6.3	6.3	2.8	2.5	2.7	2.8	2.8	2.8
PI 321006, USDA 2001	5.5	5.2	5.3	2.3	2.1	2.2	2.3	2.5	2.4
PI 321007, USDA 2001	5.6	5.5	5.5	3.4	3.3	3.4	2.8	2.8	2.8
PI 321008, USDA 2001	5.1	5.3	5.2	2.9	2.7	2.8	2.0	2.3	2.1
PI 390261, USDA 2001	5.3	5.0	5.2	2.1	1.4	1.7	2.5	1.5	2.0
PI 390262, USDA 2001	6.1	5.8	6.0	3.0	2.3	2.6	3.0	3.0	3.0
PI 401732, USDA 2001	5.8	5.7	5.7	2.8	2.6	2.7	2.5	2.5	2.5
PI 401733, USDA 2001	6.0	6.0	6.0	3.1	2.8	3.0	2.5	2.5	2.5
WI 5551, 1994	5.1	5.5	5.3	3.0	2.9	3.0	2.5	3.0	2.8

<sup>\*</sup>Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

Cultivar (four fruits per replicate),	P. capsici OP97								
incubated 3 days at room temperature	Lesion diam. (cm)		Sporulation diam. (cm)			Sporulation density**			
tomportuato	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave	Rep 1	Rep 2	Ave
Discover M, Asgrow 2000	4.0	3.6	3.8	1.6	0.3	0.9	1.0	0.3	0.6
PI 197085, USDA 2001	2.6	2.4	2.5	1.7	1.8	1.7	0.5	1.0	0.8
PI 197088, USDA 2001	3.0	2.6	2.8	1.3	1.2	1.3	0.3	1.0	0.6
PI 249562, USDA 2001	3.2	3.5	3.4	1.1	1.2	1.2	1.0	0.5	0.8
PI 271326, USDA 2001	3.3	3.2	3.3	1.2	1.4	1.3	0.5	0.8	0.6
PI 271327, USDA 2001	3.2	3.4	3.3	1.3	1.6	1.5	1.8	1.8	1.8
WI 1983 G, 1997	3.3	3.3	3.3	1.9	1.4	1.6	2.0	1.3	1.6
WI 6632 E, 1997	3.2	3.6	3.4	1.7	1.4	1.5	1.3	1.0	1.1
WI 5207, 2000	3.5	3.6	3.5	1.5	1.7	1.6	1.5	0.8	1.1
(WI) SMR 18, 2000	3.7	3.5	3.6	1.7	1.3	1.5	1.3	1.0	1.1
(WI) GY 14, 1998	3.3	3.9	3.6	1.6	1.2	1.4	1.0	0.5	0.8

<sup>\*</sup>Experiment incubated three days due to contamination.

\*\*Sporulation Density rated on a scale of 0 to 3 where 0=none, 1=faint, 2=moderate, 3=heavy sporulation.

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#### Worksheet 3-A(15)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Resistant Cultivars	Study: Evaluation of fungicides and host resistance for
	control of Phytophthora crown rot of summer
	squash, 1999.
Section I. Initial Screening on Technic	al Feasibility of Alternatives
•	
1. Are there any location-specific restrictions that inhibit	it the use of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming count	try
1d. Other (Please describe)	·

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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# Worksheet 3-A(15)(c). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

## Section II. Existing Research Studies on Alternatives to Methyl Bromide No X Yes 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) G.J. Holmes M.E. Lancaster F.J. Louws Fungicide and Nematicide Tests, 2000 3. Publication and Date of Publication Hendersonville, North Carolina 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. **Resistant Cultivars** No X 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Using a resistant cultivar was not commercially acceptable, because there was nearly 40% of the plants killed by Phytophthora capsici with this control measure. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? These results apply to Michigan growers, who would expect to experience similar losses.

OMB Control # 2060-0482

G. J. Holmes<sup>I</sup>, M. E. Lancaster<sup>2</sup> and F. J. Louws<sup>1</sup>, <sup>1</sup>Dept Plant Pathology, NC State Univ, Box 7616, Raleigh, NC 27695, and <sup>2</sup>NC Coop. Ext. Service, Hendersonville, NC 28792

EVALUATION OF FUNGICIDES AND HOST RESISTANCE FOR CONTROL OF PHYTOPHTHORA CROWN ROT OF SUMMER SQUASH, 1999: The experiment was conducted in a commercial squash field near Hendersonville, NC (GPS coordinates: N35°19.078′; W082°25.178′) where a severe outbreak of Phytophthora crown rot (PCR) was observed on squash in the summer of 1997. The field was planted to sweet corn in 1997. Soil type was a Codorus loam. The PCR-resistant squash variety was a gray zucchini (SSXP210; Harris Moran Seeds). Treatments were randomized in four complete blocks. Plots were two rows on 4-ft centers, 20 ft long and separated by two rows of pepper. Preplant incorporated (PPI) treatments were applied immediately prior to planting using a CO<sub>2</sub> backpack sprayer equipped with a single nozzle, handheld boom, hollow cone nozzle tip (TXVK-8) and operating at 40 psi. Squash was direct seeded on 7 Jun into ridged beds. Foliar treatments were applied using the same apparatus, by a single pass on each side of the plant bed for a total volume of 56 gal/A. Ridomil Gold + Dithane foliar applications were made on 23 Jun, 7 and 20 Jul and 4 Aug. All other treatments except those that were preplant incorporated (PPI) were applied weekly beginning 16 Jun and ending 11 Aug for a total of 9 applications.

Disease incidence (% mortality) evaluations began 7 days following the first observation of disease (approximately 4 wilted plants in entire test site) when plants had approximately 6 true leaves. Disease progressed rapidly with 46% of individual plots showing greater than 80% plant death on 14 Jul. Yield was not evaluated since the main effect of the disease was plant death. Highly significant block and treatment effects were detected (P=0.01) at each evaluation and for Area Under Disease Progress Curve (AUDPC). The Ridomil Gold EC treatment provided superior control of the disease compared with all other chemical treatments. Because disease attacked early, we believe that most of the effect was due to the PPI treatment rather than subsequent foliar applications. The resistant variety also held up relatively well under the intense disease pressure. However, this variety does not possess good marketable characteristics. A postharvest evaluation (10 fruit from each treatment stored for 20 days at room temperature) yielded 2 out of 120 fruit rotting due to *P. capsici*. Louws et al. report results of the parallel study on pepper in this volume.

	Mortality (%) 1								
Product and amount/A	29 Jun	07 Jul	14 Jul	20 Jul	28 Jul	03 Aug	11 Aug	17 Aug	AUDPC
Acrobat 50WP, 0.4 lb + Dithane 75DF, 2 lb	11.7 abc	47.7 b	56.2 b	73.0 ab	71.8 b	78.5 b	81.1 b	83.0 b	3278 b
Acrobat 50WP, 0.4 lb + Kocide 2000, 1.9 lb	16.9 bc	56.7 b	66.0 b	79.3 abc	86.7 ab	91.8 abc	93.8 bc	94.5 ab	3851 bc
Acrobat 50WP, 0.4 lb + Dithane 75DF, 2 lb (stem base alt.) 2	15.6 bc	52.3 b	59.8 b	71.5 b	75.8 b	80.9 bc	82.5 b	84.1 b	3441 bc
Dithane 75DF, 2 lb	22.2 c	58.6 b	64.2 b	81.2 abc	84.1 bc	89.3 abc	88.9 bc	89.7 bc	3862 bc
Ridomil Gold 4EC, 2 pt (PPI) <sup>3</sup> , Ridomil Gold 4EC, 2 pt + Díthane 75DF, 2 lb (foliar)	0.7 a	1.9 a	3.3 a	5.6 a	9.8 a	18.4 a	25.6 a	. 29.4 a	551 a
4 Quadris 2.08F, 2 pt (PPI)	9.1 abc	61.3 b	73.1 b	86.7 abc	91.1 bc	97.1 bc	97.8 bc	98.5 bc	3974 bc
<sup>4</sup> Flint 50WG, 1 lb (PPI)	13.3 abc	68.6 b	74.3 b	91.8 bc	95.9 c	99.2 c	100 c	100 с	4204 bc
4 Sovran 50WG, 1 lb (PPI)	18.3 bc	68.5 b	80.2 Ъ	88.6 abc	-91.7 bc	95.4 abc	96.6 bc	97.0 bc	4207 bc
<sup>4</sup> Acrobat 50WP, 4 lb (PPI)	12.1 c	72.0 Ъ	81.2 b	86.6 abc	89.3 bc	93.7 abc	94.9 bc	94.9 bc	4235 bc
Resistant variety, no fungicide	0.0 a	6.9 a	11.2 a	22.1 a	23.7 a	28.5 a	30.6 a	36.5 a	1204 a
Resistant variety + Acrobat 50WP, 0.4 lb +	,								
Dithane 75DF, 2 lb	5.3 ab	7.6 a	18.7 a	21.0 a	26.9 a	27.5 a	31.8 a	34.6 a	1112 a
Non-treated	19.9 c	61.2 b	74.8 b	94.9 c	98.3 c	100 c	100 c	100 c	4295 c
LSD (P=0.05)	14.1	25.7	26.6	20.0	20.0	17.8	17.5	15.8	980

Values are the means of 4 replicate plots. Treatments followed by the same letter within a column are not significantly different (K=100, Duncan-Waller K-ratio test).

F&N Tests 55:260

<sup>&</sup>lt;sup>2</sup> alternated between spray directed at the base of the plant and directed at the entire plant (not possible until plant height was >1.5 ft; approximately 20 Jun).

<sup>&</sup>lt;sup>3</sup> PPI = preplant incorporated in top 4 inches of soil.

<sup>&</sup>lt;sup>4</sup> Followed by foliar treatment at 14 and 28 days after planting.

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#### Worksheet 3-A(16). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

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For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Soilless Culture	Study: <u>UNEP 1988, B-83, B-282, B-44</u>
-------------------------------	--

#### Section I. Initial Screening on Technical Feasibility of Alternatives

1. Are there any location-specific restriction	ns that inhibit the use of this alternative on you	ur site?
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in cor	nsuming country	
1d. Other (Please describe)	<u> </u>	
		[2] 6. (元)

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#### Worksheet 3-A(16). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes X No\_\_\_\_ 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Soilless Culture 6. Was crop yield measured in the study? Yes \_\_\_\_\_ No 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? Soilless culture is achieved in some parts of the world through the use of volcanic gravel, and has been helpful in managing various soil-borne pathogens. This method of disease control is unproven for management of Phytophthora capsici, and is not feasible for Michigan growers who do not have access to volcanic gravel.

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#### Worksheet 3-A(17). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

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When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### BACKGROUND

1. A

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Substrates, Plug Plants	Study: UNEP 1988, B-83, B-90, B-94, B-282	

#### Section I. Initial Screening on Technical Feasibility of Alternatives

a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming country		
1d. Other (Please describe)		
1d. Other (Please describe)		

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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ID#	

#### Worksheet 3-A(17). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? Yes X No 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) 3. Publication and Date of Publication 4. Location of research study 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Substrates, Plug Plants 6. Was crop yield measured in the study? Yes \_\_\_\_ 7. Describe the effectiveness of the alternative in controlling pests in the study. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of these studies do not apply to the situation in Michigan, because Phytophthora capsici is not disseminated via seeds or transplants. The examples given in the UNEP 1998 studies include Alternaria. Didymella, Fusarium oxysporum, Clavibacter michiganensis subsp. michiganensis, Verticillium spp. and

Pseudomonas spp. These examples do not apply to the situation in Michigan. Use of pathogen-free seeds and

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transplants is not a viable alternative for P. capsici.

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			ID:

#### Worksheet 3-A(18). Alternatives - Technical Feasibility of Alternatives to Methyl Bromue

In this worksheet, you should address why an alternative pest management strategy on the list (see previous page) is or is not effective for your conditions. This worksheet contains 9 questions. You must complete one copy of worksheet 3-A for each research study you use to evaluate a single methyl bromide alternative. Use additional pages as need.

For worksheet 3-A you must complete one worksheet for each alternative, for each research study addressed. Please number the worksheets as follows. For the same alternative, first research study, label the worksheet 3-A(1)(a). For the same alternative, second research study, label the worksheet 3-A(1)(b). For the first alternative, third research study, label the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)(a). For the second alternative, second research study, label the worksheet 3-(A)(2)(b).

When completing Section II, if you cite a study that is on the EPA website, you only need to complete questions 1, 5, and 8.

Summarize each of the research studies you cite in the Research Summary Worksheet.

If you prefer, you may provide the information requested in this worksheet in a narrative review of one or more relevant research reports. The narrative review must reply to Section I and questions 1 through 8 in Section II. A Research Summary Worksheet of relevant treatments should be provided for each study reviewed.

#### **BACKGROUND**

EPA must consider whether alternative pest control measures (pesticide and non-pesticidal, and their combination) could be used -successfully instead of methyl bromide by crop and circumstance (geographic area.) The Agency has developed a list of possible alternative pest control regimens for various crops, which can be found at http://www.epa.gov/ozone/mbr or by calling 1-800-296-1996.

There are three major ways you can provide the Agency with proof of your investigative work.

- (1) Conduct and submit your own research
- (2) Cite research that has been conducted by others
- (3) Cite research listed on the EPA website

Whether you conduct the research yourself or cite studies developed by others, it is important that the studies be conducted in a scientifically sound manner. The studies should include a description of the experimental methodology used, such as application rates, application intervals, pest pressure, weather conditions, varieties of the crop used, etc. All results should be included, regardless of outcome. You must submit copies of each study to EPA unless they are listed on the Agency website.

The Agency has posted many research studies on a variety of crops on its website and knows of more studies currently in progress. EPA will add studies to its website as they become publicly available. You are encouraged to review the EPA website and other websites for studies that pertain to your crop and geographic area.

In addition, EPA acknowledges that, for certain circumstances, some alternatives are not technically feasible and therefore no research has been conducted (i.e. solarization may not be feasible in Seattle). You should look at the list of alternatives provided by the Agency and explain why they cannot be used for your crop and in your geographic area.

Use additional pages as needed.

Alternative: Fungicides .	Study: The dynamics of mefenoxam insensitivity in a	а
	recombining population of Phytophthora cap	sici
	characterized with amplified fragment length	
	polymorphism markers.	
Section I. Initial Screening on Technic	al Feasibility of Alternatives	
_	•	
1. Are there any location-specific restrictions that inhibi	the use of this alternative on your site?	
1a. Full use permitted	X	
1b. Township caps		
1c. Alternative not acceptable in consuming count	y	
1d. Other (Please describe)		

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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ID#	

## Worksheet 3-A(18). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

<ol> <li>Is the study on EPA's website</li> </ol>	? Yes	NoX
1a. If not on the EPA w	ebsite, please attach a copy.	
2. Author(s) or researcher(s)	K.H. Lamour	
	M.K. Hausbeck	
. Publication and Date of Public	Phytopathology 91:5	53-557, 2002
. Location of research study	Michigan, USA	
. Name of alternative(s) in stud	/. If more than one alternative, list	•
. Was crop yield measured in th	e study? Yes	NoX
		ed fungicide, is common in Michigan fields.
Insensitivity of the pathogen to the	nis fungicide renders this treatment	ineffective.
. Discuss how the results of the other factors that would affect		ould you expect similar results? Are there

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# The Dynamics of Mefenoxam Insensitivity in a Recombining Population of *Phytophthora capsici* Characterized with Amplified Fragment Length Polymorphism Markers

K. H. Lamour and M. K. Hausbeck

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#### **ABSTRACT**

Lamour, K. H., and Hausbeck, M. K. 2001. The dynamics of mefenoxam insensitivity in a recombining population of *Phytophthora capsici* characterized with amplified fragment length polymorphism markers. Phytopathology 91:553-557.

Recent findings from Michigan suggest that recombination may play a role in the survival and evolution of sensitivity to the fungicide mefenoxam in populations of *Phytophthora capsici* on cucurbit hosts. In 1998, 63 mefenoxam insensitive isolates were recovered from a squash field in which mefenoxam had been applied. Additional isolates were recovered from untreated squash fields planted at this location in 1999 (200 isolates) and the spring of 2000 (34 isolates). Isolates from 1998 and 1999 were characterized using fluorescent amplified fragment length polymorphism (AFLP) markers and all isolates were screened for com-

patibility type and mefenoxam sensitivity. In 1998 and 1999, 92 and 71% of the isolates, respectively, had unique multilocus AFLP genotypes with no identical isolates recovered between years. Seventy-two identical AFLP markers were clearly resolved in both the 1998 and 1999 sample sets, and fixation indices for the 37 polymorphic AFLP loci indicate little differentiation between years. There was no decrease in the frequency of resistant isolates during the 2 years without mefenoxam selection. We conclude that oospores play a key role in overwintering and that the frequency of mefenoxam insensitivity may not decrease in an agriculturally significant time period (2 years) once mefenoxam selection pressure is removed.

Additional keywords: fungicide resistance, genetic diversity, population genetics.

Crown, root, and fruit rot caused by Phytophthora capsici is increasing in Michigan cucurbit production fields, and uninfested land suitable for rotation is becoming increasingly scarce, especially in areas undergoing rapid urban development. The phenylamide fungicide (PAF) mefenoxam is a systemic fungicide that appears to be acting at the level of DNA translation, and is fungistatic to fully sensitive isolates of P. capsici (2,13). Although mefenoxam has been considered by some growers to be helpful, mefenoxam insensitive isolates were reported on bell peppers in North Carolina and New Jersey by Parra and Ristaino in 1998 (18) and have since been recovered from 10 of 11 farms sampled in Michigan (13), as well as, in Georgia (15) and southern Italy (19). Mefenoxam insensitivity in Michigan P. capsici isolates is inherited as a single gene exhibiting incomplete dominance (13), which is consistent with the reports for a variety of other oomycetous organisms (2). Investigations with P. infestans indicate that insensitivity may be conferred by genes at different chromosomal positions (5), suggesting that the basis of insensitivity in different populations may not be identical. Sexual recombination, in particular, has the potential to impact management strategies that employ PAFs because the fully insensitive (two copies of the insensitivity allele) phenotype may be directly generated. P. capsici is heterothallic and the sexual stage is initiated when isolates of opposite compatibility type, designated A1 and A2, come into close association to form thick-walled oospores (4). The asexual stage includes the production of caducous sporangia born on long pedicels, which may release motile zoospores if free water is present. Asexual spores are thought to be responsible for the polyyclic nature of disease development (20).

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PAF resistance in the genus Phytophthora and, in particular, the P. infestans-potato pathosystem, is well documented (2,4,9). Until recently, the population structure of P. infestans appeared to be largely clonal outside of P. infestans putative center of origin (6). The recent detection of both P. infestans compatibility types along with increased genotypic diversity in some potato growing regions indicates that the sexual stage is likely active and may significantly impact control strategies that have proved useful in the past (3,8). When PAF resistance in European P. infestans populations increased significantly in the early 1980s, the efficacy of the PAF metalaxyl was only regained after the product was not made available to growers for a period of time (2). This strategy apparently allowed the resistant populations to decline or become extinct and depends on ephemeral populations or, in the case of resident populations, upon a significant cost for resistance outside of selection pressure. A recent study of sensitive versus PAF resistant P. nicotianae isolates from citrus suggests negligible fitness costs for PAF resistance and reports that 2 years without PAF use did not reduce the proportion of resistant isolates in groves (21). Kadish and Cohen report that PAF-resistant P. infestans isolates in Israel were more aggressive in colonizing tuber tissue than sensitive isolates (12).

Novel techniques have been developed recently that allow characterization of DNA-level polymorphism in organisms for which little is known about the genome. An example is the amplified fragment length polymorphism (AFLP) technique introduced by Vos et al. in 1995 (23). This technique relies on restriction enzyme fragmentation of genomic DNA with the concomitant ligation of synthetic adaptors to the DNA fragment ends. Stringent polymerase chain reaction (PCR) amplification using adaptor-complementary primers with additional selective nucleotides allow for the amplification of fragment subsets. DNA fragment subsets are termed fingerprints and may be resolved with a range of techniques (1). AFLP markers have been used on a variety of organ-

isms (14,22) and the procedure generates a large number of reproducible markers (1,22). The limitation that these markers are generally scored as dominant markers (e.g., either present or absent) for diploid organisms requires the use of relatively large sample sets (11,25).

Our null hypotheses are that sexual recombination has a significant impact on the population structure of *P. capsici* in Michigan and that mefenoxam insensitivity may not decrease in the time frame of a typical 2-year rotation outside of mefenoxam selection pressure.

#### MATERIALS AND METHODS

Field plot. Research was conducted on a commercial farm in southwest Michigan, with a history (>11 years) of *P. capsici* on bell peppers and squash and intensive use of PAF. The 4.05-ha field sampled had previously been cropped to soybeans and corn with no known record of *P. capsici* susceptible crops (e.g., tomatoes, peppers, or cucurbits) prior to 1997. During 1997 and 1998, yellow squash and zucchini grown in this field became diseased with *Phytophthora* crown, root, and fruit rot and the grower applied mefenoxam as part of a disease management strategy (Novartis, Greensboro, NC). In 1998, all isolates recovered were either intermediately or fully insensitive to mefenoxam. Both A1

TABLE 1. Fixation indices  $(F_{ST})$  for 37 amplified fragment length polymorphism loci from unique *Phytophthora capsici* isolates collected from a single Michigan cucurbit field during 1998 (N = 57) and 1999 (N = 141)

Fragment <sup>a</sup>	1998 f(aa) <sup>b</sup>	1999 f(aa)	$F_{\mathrm{ST}}^{\mathrm{c}}$	
45	0.02	0.06	0.018	
54	0.29	0.29	0.000	
64	0.82	0.55	0.048	
104	0.11	0.06	0.007	
106	0.11	0.04	0.025	
110	0.41	0.36	0.002	
130	0.41	0.30	0.009	
146	0.47	0.24	0.038	
149	0.12	0.27	0.029	
154	0.39	0.31	0.004	
156	0.53	0.83	0.054	
172	0.56	0.33	0.034	
189	0.16	0.56	0.121	
192	0.16	0.37	0.044	
193	0.35	0.20	0.022	
211	0.47	0.15	0.088	
241	0.48	0.32	0.018	
256	0.04	0.01	0.022	
258	0.43	0.49	0.002	
261	0.55	0.54	0.000	
270	0.57	0.41	0.015	
282	0.35	0.40	0.002	
285	0.51	0.73	0.030	
314	0.51	0.34	0.019	
320	0.41	0.51	0.006	
333	0.16	0.20	0.002	
346	0.36	0.33	0.001	
361	0.33	0.49	0.017	
383	0.21	0.15	0.005	
418	0.40	0.34	0.002	
431	0.34	0.32	0.001	
438	0.67	0.45	0.028	
454	0.65	0.49	0.015	
492	0.29	0.40	0.009	
504	0.51	0.47	0.001	
511	0.38	0.28	0.007	
548	0.78	0.78	0.000	

<sup>&</sup>lt;sup>a</sup> EcoRI-AC/MseI-CA selectively amplified fragment size in base pairs.

and A2 compatibility types were present, and oospores were detected in diseased fruit. In 1999 and 2000, yellow squash was established in a 1,124-m<sup>2</sup> experimental plot in this field, and mefenoxam was not applied. Diseased plants and fruit were sampled on 20 August 1998 (63 isolates from entire field), June through August 1999 (200 isolates from experimental plot), and 13 July 2000 (34 isolates from experimental plot). All isolates were recovered from single diseased plants or fruit.

Isolate collection and maintenance. Isolation from diseased plant material was made onto BARP (25 ppm of benomyl, 100 ppm of ampicillin, 30 ppm of rifampicin, and 100 ppm of pentachloronitrobenzene)-amended UCV8 (840 ml of distilled water, 163 ml of unclarified V8 juice, 3 g of CaCO<sub>3</sub>, and 16 g of Bacto agar) plates. Procedures for obtaining single zoospore isolates were as previously described (13). Single zoospore cultures were maintained on 30 ppm of rifampicin and 100 ppm of ampicillin (RA)-UCV8 plates and transferred bimonthly. Long-term storage consisted of a single 7-mm plug of expanding mycelium from each single zoospore culture being placed in a 1.5-ml microfuge tube with one sterilized hemp seed and 1 ml of sterile distilled water, incubated for 2 to 3 weeks at 23 to 25°C, and stored at 15°C long term.

Phenotypic characterization. Isolates were screened for compatibility type as previously described (13). Mefenoxam sensitivity was characterized according to the in vitro screening technique described by Lamour and Hausbeck (LH technique) for P. capsici isolates in Michigan (13). Isolates were scored as sensitive (S) if growth on UC-V8 agar amended with 100 ppm of mefenoxam was less than 30% compared with a control, as intermediately sensitive (IS) if between 30 and 90%, and fully insensitive (I) if greater than 90% compared with the unamended control. These mefenoxam sensitivity categories are based on a trimodal distribution of 523 field isolates of P. capsici. Clear modal distributions were only attained when screening was conducted with a single high rate of mefenoxam-amended (100 ppm) media (K. Lamour, unpublished data). These putative mefenoxam sensitivity categories were tested by in vitro crosses (I  $\times$  S, IS  $\times$  IS, IS  $\times$  S, and S x S), and chi-square analysis confirmed that the observed progeny numbers were not significantly different than expected for Mendelian inheritance of an incompletely dominant trait (13).

The LH technique differs from a commonly used method described by Goodwin, Sujkowski, and Fry (GSF technique) (9) for *P. infestans* which uses two levels of amended media (5 and 100 ppm) to differentiate the three mefenoxam sensitivity phenotypes and which has been used to characterize *P. capsici* isolates (15,18,19). Unfortunately, analysis of our in vitro crosses and field isolates by the GSF technique did not resolve a clear modal distribution (K. Lamour, *unpublished data*). Assignment of Michigan *P. capsici* isolates to the S category was the same whether using the LH or GSF technique. The only difference was that some *P. capsici* isolates from Michigan rated as fully insensitive by the GSF technique were rated as intermediately sensitive by the LH technique.

DNA extraction and AFLP fingerprinting. A technique for avoiding bacterial contamination prior to growing isolates for DNA extraction was implemented using a modified Van Teigham cell (4). The uppermost portion of a 7-mm plug of mycelium was placed onto the surface of RA-WA plates (30 ppm of rifampicin, 100 ppm of ampicillin, 1,000 ml of distilled water, and 16 g of Bacto agar) and an autoclaved cap from a 1.5-ml microfuge tube was placed over the plug which forced the isolate to grow through the amended media. Isolates were incubated in the dark for 2 t 3 days before two 7-mm plugs were transferred to approximately 15 ml of RA-UCV8 broth in petri dishes (100 × 15 mm) and incubated in the dark for 3 days at 23 to 25°C. Mycelial mats were washed with distilled water and dried briefly under vacuum before being frozen to -20°C and lyophilized.

b Observed frequency of the absent state where "a" represents the absence of a fragment.

c F<sub>ST</sub> calculated from estimated allele frequencies. According to Wright's qualitative guidelines, values from 0 to 0.05 indicate little genetic differentiation and values from 0.05 to 0.15 indicate moderate genetic differentiation.

Lyophilized mats were ground with a sterile mortar and pestle. Whole genomic DNA from approximately 50 mg of ground mycelium was extracted with a plant mini kit (Dneasy: Oiagen Inc., Valencia, CA) according to the manufacturers directions. DNA was quantified (Nucleic Acid QuickSticks; Clontech, Palo Alto, CA) according to the manufacturers directions and approximately 100 ng of DNA was subjected to a restriction/ligation reaction, preselective amplification, and selective amplifications using the PCR core mix, adaptor sequences, core primer sequences, and fluorescent-labeled primers available in an AFLP microbial fingerprinting kit (Perkin-Elmer Applied Biosystems, Foster City, CA, henceforth referred to as PE/ABI) and performed exactly as described in protocol part 402977 Rev A (23). All PCR reactions were performed with a minicycler (MJ Research Inc., Waltham, MA) in 0.2-ml tubes according to the cycling parameters outlined in the microbial fingerprinting protocol.

An initial optimization set of reactions was performed with preselective products from *P. capsici* isolate OP97, which was isolated from a cucumber fruit in 1997 (13). Selective amplifications with the selective primers *EcoRI*-AA, -AC, -AG, and -AT were performed in all 16 combinations with the *MseI*-CA, -CC, -CG, and -CT selective primers. *EcoRI* selective primers, available from PE/ABI, were labeled at the 5' end with either carboxy-fluorescein (FAM), carboxytetramethyrhodamine (TAMRA), or carboxy-4',5'-dichloro-2',7'-dimethoxyfluorescein (JOE) fluorescent dyes. The fluorescent dyes are excited by laser radiation and visualized by their characteristic absorption-emission frequencies. Only the fragments\_containing an *EcoRI* restriction site are resolved.

Products from three reactions labeled with different colored dyes and a carboxy-X-rhodamine (ROX) size standard were loaded into each lane on a denaturing polyacrylamide gel and the fragments resolved in a DNA sequencer (ABI Prism 377). Results were prepared for analysis in the form of electropherograms using GeneScan Analysis software (PE/ABI). AFLP fragments were scored manually as present (1) or absent (0) using Genotyper (PE/ABI). Only DNA bands that consistently exhibited unambiguous presence or absence profiles were scored.

A single isolate, OP97, was subjected to the aforementioned protocol using three primer pair combinations that were chosen as optimal on three separate occasions, approximately 3 months apart, to test for reproducibility of AFLP profiles.

Clone detection and cluster analysis. AFLP fragments were considered polymorphic if the most common allele was present in less than 95% of the isolates from a given sample set and scored for presence (1) or absence (0) (10). AFLP fragments present in more than 95% of the isolates from a given sample set were considered monomorphic. Analysis of the resulting binary data matrix was performed using NTSYS-pc version 2.02k (Exeter Software, Setauket, NY). Unweighted pair group method with arithmetic averages cluster analysis was performed on the matrix of similarity coefficients calculated from all possible pairwise comparisons of individuals within and among the 1998 and 1999 populations and a tree generated. Isolates showing complete homology at all loci were considered to be clones and except for a single representative isolate were excluded from frequency calculations.

Allele frequency and fixation indices. Allele frequencies for AFLP markers were estimated utilizing the expected relationship between gene and genotype frequencies in a randomly mating population (i.e., Hardy-Weinberg proportions). The frequency of the recessive (absent) allele (q) was calculated from the observed number of recessive homozygote individuals (X) in a sample of n individuals by the formula for dominant markers described by Jorde et al. (11):

$$\hat{q} = \sqrt{x + \frac{1 - x}{4n}}$$

where x = X/n is the observed proportion of individuals that do not display the dominant (present) marker phenotype. In order to test whether the composite genetic profiles from 1998 and 1999 were consistent with a single randomly mating population, the fixation index was calculated for each AFLP loci from the variance in allele frequencies according to the following formula:  $F_{ST} = [(p_1 - p_2)^2/4]/(average p \times average q)$ , where p is the allele frequency for the present state with  $p_1$  and  $p_2$  indicating the two sample populations, and q is the allele frequency for the absent state (10). Fixation indices for individual loci were interpreted according to the qualitative guidelines suggested by Wright (24), where the range 0 to 0.05 indicates little genetic differentiation, range 0.05 to 0.15 indicates moderate genetic differentiation, and greater than 0.25 indicates great genetic differentiation (10).

#### RESULTS

AFLP band characterization. Evaluation of the 16 EcoRI + 2-MseI + 2 selective primer pair combinations indicated that EcoRI + AC-MseI + CA gave the most clearly resolved fragment profile and was used to amplify genomic DNA from all isolates in both the 1998 and 1999 sample sets. This primer combination resulted in 72 clearly resolved fragments of which 37 (51%) fragments were polymorphic in both 1998 and 1999 (Table 1). All 72 fragments were present in both 1998 and 1999 and no novel fragments were detected between years. The following 35 fragments (size in base pairs) were monomorphic in both the 1998 and 1999 sample sets: 41, 43, 47, 49, 58, 66, 70, 82, 85, 114, 118, 123, 133. 135, 140, 159, 174, 235, 247, 249, 272, 278, 295, 298, 300, 341, 351, 355, 367, 402, 474, 488, 502, 519, and 527. AFLP profiles for isolate OP97, generated from separate DNA extractions on three separate occasions over a 1-year period, resulted in identical banding patterns with the only difference being minor changes in the intensity of the electropherogram signal. Occasionally individual reactions resulted in poorly resolved fingerprint profiles (e.g., low intensity of signal) and were repeated until signals were deemed optimal.

Phenotypic, genotypic, and gene diversity. No isolates sensitive to mefenoxam were recovered in 1998 or 2000, and single A1 sensitive and A2 sensitive isolates were recovered in 1999 (Table 2). In 1998, 18% of the isolates were intermediately sensitive and 82% were insensitive, in 1999, 2% were sensitive, 28% were intermediately sensitive and 70% were insensitive, and in 2000, 15% of the isolates were intermediately sensitive and 85% were insensitive to mefenoxam (Table 2).

In 1998, 57 of the 63 isolates recovered, and 141 of the 200 isolates recovered in 1999 were unique based on multilocus AFLP profiles. No identical multilocus genotypes were recovered between 1998 and 1999. Five isolates (two A2/I, two A2/IS, and

TABLE 2. Phenotypic diversity of *Phytophthora capsici* isolates recovered from the same cucurbit field in 1998, 1999, and 2000

	No. of	Compatibility type and mefenoxam sensitivi				rityc	
Yeara	isolates <sup>b</sup>	A1/S	A1/IS	A1/I	A2/S	A2/IS	A2/I
1998	57	_	4	31	_	6	16
1999	141	1(2)	17 (20)	57 (53)	1(1)	23 (18)	42 (47)
2000	34	- '	2	8 `	- ` ´	3 ` ´	21 ` ´

<sup>&</sup>lt;sup>a</sup> Mefenoxam was applied in 1998 but not in 1999 or 2000.

b Sample sets from 1998 and 1999 consist of unique multilocus genotypes as determined with amplified fragment length polymorphism fingerprinting. The 2000 sample set was recovered at the beginning of the growing season and was not fingerprinted.

<sup>&</sup>lt;sup>c</sup> S = sensitive, IS = intermediately sensitive, and I = insensitive as determined by in vitro screening on 100 ppm of mefenoxam-amended agar. Numbers in parentheses indicate the expected number of isolates when mefenoxam insensitivity is assumed to be controlled by a single incompletely dominant gene in Hardy-Weinberg equilibrium unlinked to compatibility type.

one A1/I) of *P. capsici* collected in 1998 had one clonal representative. Fourteen isolates collected in 1999 had between two and four clones (Table 3). A single A1 compatibility type insensitive isolate had 40 clones recovered over the course of the 1999 season and comprised 3% of the early, 15% of the mid-, and 43% of the late sampling intervals (Table 3). The 1999 sampling intervals (early, mid, and late) are based on the dates of sampling and are not intended to reflect stages of plant growth or the epidemiology of *P. capsici*. Cluster analysis of AFLP fingerprint variation indicated no significant clustering of isolates between 1998 and 1999.

The majority (98%) of the 37 polymorphic AFLP fragments showed little genetic differentiation ( $F_{\rm ST}$  < 0.05) between 1998 and 1999 according to Wrights qualitative criterion (Table 1) (24).

#### DISCUSSION

P. capsici causes significant damage to cucurbit hosts in Michigan each year. In an effort to prevent or control epidemics, many growers have used either metalaxyl or the newer, but similarly acting compound, mefenoxam as a part of their disease management strategy. This study was initiated in an effort to address the concerns of growers who have high levels of mefenoxam insensitivity.

Phenotypic data (mefenoxam sensitivity and compatibility type) from a 1998 survey suggested that insensitivity to mefenoxam was common and that some level of recombination is occurring in the field (13), but without the application of additional polymorphic markers our ability to assess population structure was severely restricted. AFLP analysis proved to be a powerful tool for resolving the population dynamics of *P. capsici*. A single selective primer combination, *EcoRI-AC-MseI-CA*, generated 72 bands of which 37 were polymorphic in our 1998 and 1999 sample sets. AFLP fingerprinting, in conjunction with temporal sampling, provided a useful characterization of *P. capsici* from one season to the next and allowed us to track asexual disease development over the course of a single season.

Our data suggests that sexual recombination significantly impacts the structure of this *P. capsici* population. The finding that 198 of the 262 isolates recovered between 1998 and 1999 had unique multilocus AFLP genotypes is consistent with the high level of genotypic diversity expected in an outcrossing population

TABLE 3. Clone contribution of 15 *Phytophthora capsici* isolates to the total number of isolates collected in 1999 (N = 200)

			` ,		
No. of clones in early, mid, and late season					nd late season <sup>c</sup>
Isolate	No. of clones <sup>a</sup>	CT/MSb	6/22 - 7/16 N = 60	7/20 - 8/3 N = 80	8/5 - 8/18 N = 60
JP571	2	A1/I	2	_	_
JP583	2	A1/I	2	-	
JP944	3	A1/I	2	1	
JP999	3	A1/I	2	1	-
JP1007	2	A1/I	1	1	_
JP1042	2	A2/I	1	1	_
JP1096	2	A1/I	_	1	1
JP1102	2	A2/I	_	2	
JP1215	3	A2/I	3	_	_
JP1342	2	A2/IS	_	2	
JP1369	2	A1/I	1	1	_
JP1384	4	A2/I	3	1	_
JP1512	2	A1/I	1	_	1
JP1555	3	A1/I	_		3
JP1632	40	A1/I	2	12	26

<sup>&</sup>lt;sup>a</sup> Total number of isolates with identical multilocus amplified fragment length polymorphism profiles.

<sup>c</sup> Sample intervals based on sampling dates only.

(7,16,17). Although clonal reproduction occurred in 1998 and 1999, no identical genotypes were recovered between years, suggesting that oospores are important for overwintering. The finding that 35 of the 37 polymorphic fragments exhibited very little differentiation (i.e., change in allele frequency) based on the estimated fixation indices between 1998 and 1999 is consistent with the expectations for a recombining population large enough to avoid dramatic changes due to genetic drift.

In 1999 and 2000, sensitive and intermediately sensitive isolates (42 of 175) did not increase in a manner suggesting selection in favor of mefenoxam sensitivity outside of mefenoxam selection pressure. The fact that 14 of the 15 isolates with clonal reproduction in 1999 were fully insensitive may be another indication that mefenoxam insensitivity does not have significant costs outside of mefenoxam selection pressure. If we assume that there is only a single mefenoxam insensitivity gene in this population unlinked to compatibility type, designated I, and that this population is effectively free from the effects of migration and genetic drift, some interesting speculations can be made. For instance, in 1999, if the mefenoxam sensitivity phenotypes are assumed to represent genotypes (e.g., a fully insensitive isolate has two copies of the I allele) then the frequency of I can be estimated and the observed number of unique isolates that fall into each of the six mefenoxam sensitivity/compatibility type categories can be compared with the expectations under Hardy-Weinberg equilibrium. In 1999, the estimated frequency of I was 0.84, and chi-square analysis, using the data in Table 2, indicates that the observed numbers do not differ from those expected under Hardy-Weinberg equilibria at  $P = 0.50 \ (\chi^2 = 3.09, \ df = 4)$ . Although this is not a particularly powerful test due to the large number of assumptions (10), it does lend support to the hypothesis that this population meets the criterion for panmixia.

Our results do not allow us to reject the null hypothesis that sexual recombination significantly impacts the structure of this population. It appears that sexual recombination plays a significant role in maintaining genotypic and gene diversity while concomitantly producing overwintering inoculum. Our data also suggest that sexual recombination may serve as a potent force for integrating a beneficial allele based on the finding that there were a total of 133 unique multilocus genotypes fully insensitive to mefenoxam between 1998 and 1999. An interesting question that can only be answered by following a fully sensitive population as it shifts to insensitivity is how much genetic diversity is lost, if any, during the PAF selection process? The question of how long mefenoxam resistance will remain in a population of P. capsici when selection pressure is removed can only be answered in a tentative way. It appears that in this population, insensitivity will not decrease within the time frame of a typical 2-year rotation and, once resistance to mefenoxam is established, the future usefulness of this fungicide may be extremely limited.

Comparison of the population structure reported at this single location is currently being compared with other locations in Michigan and the United States and should provide useful insight into the amount of genetic diversity in sensitive versus insensitive populations as well as the contribution of migration to *P. capsici* population structure.

#### **ACKNOWLEDGMENTS**

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b CT = compatibility type and MS = mefenoxam sensitivity where S = sensitive, IS = intermediately sensitive, and I = insensitive as determined by in vitro screening on 100 ppm of mefenoxam-amended agar.

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Worksheet 3-A(18)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide
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In this worksheet, you should address why an alternative pest not effective for your conditions. This worksheet contains 9 q each research study you use to evaluate a single methyl brom	uestions. You must complete one copy of worksheet 3-A for
For worksheet 3-A you must complete one worksheet for each number the worksheets as follows. For the same alternative, f same alternative, second research study, label the worksheet the worksheet 3-A(1)(c). For the second alternative, first research study, label the worksheet 3-(A)(2)	First research study, label the worksheet 3-A(1)(a). For the 3-A(1)(b). For the first alternative, third research study, label trch study, label the worksheet 3-(A)(2)(a). For the second
When completing Section II, if you cite a study that is on the E	PA website, you only need to complete questions 1, 5, and 8.
Summarize each of the research studies you cite in the Resear	ch Summary Worksheet.
If you prefer, you may provide the information requested in this research reports. The narrative review must reply to Section I Worksheet of relevant treatments should be provided for each	and questions 1 through 8 in Section II. A Research Summary
BACKGROUND	
EPA must consider whether alternative pest control measures (pessuccessfully instead of methyl bromide by crop and circumstance (alternative pest control regimens for various crops, which can be for	
There are three major ways you can provide the Agency with proof (1) Conduct and submit your own research (2) Cite research that has been conducted by others (3) Cite research listed on the EPA website	of your investigative work.
Whether you conduct the research yourself or cite studies develope scientifically sound manner. The studies should include a description application intervals, pest pressure, weather conditions, varieties of outcome. You must submit copies of each study to EPA unless the study the study to EPA unless the study to EPA unless the study to EPA	on of the experimental methodology used, such as application rates, the crop used, etc. All results should be included, regardless of
The Agency has posted many research studies on a variety of crope EPA will add studies to its website as they become publicly available websites for studies that pertain to your crop and geographic area.	
In addition, EPA acknowledges that, for certain circumstances, som research has been conducted (i.e. solarization may not be feasible the Agency and explain why they cannot be used for your crop and	in Seattle). You should look at the list of alternatives provided by
Use additional pa	ges as needed.
Alternative: Fungicides Study:	The spatiotemporal genetic structure of Phytophthora capsici in Michigan and implications
	for disease management.
Section I. Initial Screening on Technical Fe	asibility of Alternatives
1. Are there any location-specific restrictions that inhibit the us	e of this alternative on your site?
1a. Full use permitted	X
1b. Township caps	
1c. Alternative not acceptable in consuming country	
1d. Other (Please describe)	

If use of this alternative is precluded by regulatory restriction for all users covered by this application, the applicant should not complete Section II.

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#### Worksheet 3-A(18)(b). Alternatives - Technical Feasibility of Alternatives to Methyl Bromide

# Section II. Existing Research Studies on Alternatives to Methyl Bromide 1. Is the study on EPA's website? 1a. If not on the EPA website, please attach a copy. 2. Author(s) or researcher(s) K.H. Lamour M.K. Hausbeck 3. Publication and Date of Publication Phytopathology 92:681-684, 2002 4. Location of research study Michigan, USA 5. Name of alternative(s) in study. If more than one alternative, list the ones you wish to discuss. Mefenoxam 6. Was crop yield measured in the study? 7. Describe the effectiveness of the alternative in controlling pests in the study. Insensitivity of *Phytophthora capsici* to mefenoxam, a commonly used fungicide, is common in Michigan fields. Insensitivity of the pathogen to this fungicide renders this treatment ineffective. 8. Discuss how the results of the study apply to your situation. Would you expect similar results? Are there other factors that would affect your adoption of this tool? The results of this study are directly applicable since the research was conducted in Michigan, USA.

OMB Control # 2060-0482

#### New Frontiers in Plant Disease Losses and Disease Management

# The Spatiotemporal Genetic Structure of *Phytophthora capsici* in Michigan and Implications for Disease Management

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Root, crown, and fruit rot caused by *Phytophthora capsici* Leonian is a limiting factor for the production of peppers, tomatoes, and cucurbit crops in Michigan and the United States. Like many species in the genus *Phytophthora*, *P. capsici* has the potential for rapid polycyclic disease development from a limited amount of initial inoculum (6). *P. capsici* produces caducous sporangia that can be spread by wind-blown rain or release 20 to 40 motile zoospores in the presence of free water. The polycyclic phase of disease development is thought to be driven primarily by asexual spore dispersal at a local scale (within and down rows). Sexual reproduction requires both the A1 and A2 compatibility types (CTs) and results in the production of thick-walled oospores. Oospores are thought to serve as the primary survival structure outside of host tissue.

Recommended disease management strategies stress the importance of avoiding excess water in the plant rhizosphere by using well-drained fields, conservative irrigation, and planting on raised beds. Additional recommendations include rotation to nonsusceptible hosts for at least 2 years and the use of fungicides. The phenylamide fungicide (PAF) mefenoxam is a systemic compound with high activity against P. capsici and has been used by growers throughout the United States to control P. capsici. Insensitivity to PAF has been reported for a number of other oomycetous organisms (Bremia lactucae, P. infestans, and P. sojae, etc.) and appears to be conferred by a single incompletely dominant gene of major effect (1). Growers in Michigan practicing 2+-year rotation in well-drained fields using an array of fungicidal management tools have experienced significant losses to P. capsici. Michigan is the number one producer of cucumbers for pickling in the United States and it was at the request of grower groups associated with this industry that research into the epidemiology and reproductive biology of P. capsici on cucurbit hosts was initiated.

Although many researchers cite oospores as the most likely propagule for survival outside of host tissue, there have been very few investigations specifically aimed at determining the impact of sexual reproduction in natural populations of *P. capsici*. Our hypothesis was that the sexual stage may play an important role not only in survival but also in the adaptation of *P. capsici* populations to environmental stresses (e.g., fungicides). Our goal was to perform a comprehensive investigation of the phenotypic and genetic diversity present in *P. capsici* populations from the major vegetable production regions of Michigan, with the implicit intention of addressing questions concerning epidemiology, repro-

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ductive biology, and the durability of currently recommended management strategies.

#### **METHODOLOGY** -

Isolate collection and maintenance. Sampling of diseased fields began at the end of the 1997 growing season and continued through September 2000. In all cases, fields were sampled on a grid with quadrants varying from 40 m² to 12 km². A limited number of isolates were collected in 1997. In 1998, the strategy was to collect as many samples from as many fields as possible. This strategy was modified in 1999 and 2000 to focus on specific fields. Isolations from diseased plants were made onto selective media and single zoospore cultures were generated according to standard single sporing techniques (3). Isolates were placed into long-term storage (15°C) using a hemp seed/sterile water technique.

Phenotypic characterization. Single zoospore isolates were screened for CT using known A1 and A2 isolates. In vitro screening techniques published for other Phytophthora species for assessing sensitivity to mefenoxam were compared and a novel, simple, high dose screen using 100 ppm of mefenoxam-amended V8 agar was found to separate field isolates into three modal distributions that appeared consistent with the expectations of a single incompletely dominant gene governing mefenoxam insensitivity (e.g., sensitive, intermediately sensitive, and fully insensitive). These putative mefenoxam sensitivity (MS) groupings were tested by performing a series of crosses and testing whether the observed progeny sets met the expectations for Mendelian inheritance of a single incompletely dominant gene controlling insensitivity to mefenoxam. Sexual crosses were conducted on unclarified V8 agar plates and incubated for 3 months in the dark. Individual germinated oospores were recovered after 3 months using previously published techniques (2).

The efficacy of this in vitro mefenoxam screening technique was further tested in pumpkin seedlings using progeny from a cross between parents intermediately sensitive to mefenoxam. Nine isolates from each of the three MS categories were screened for pathogenicity on untreated seedlings. Single sensitive, intermediately sensitive, and fully insensitive isolates were then placed onto the unwounded surface of plants treated with either a field rate of mefenoxam, three times the field rate, or distilled water. Lesion diameters on seedling stems were measured after 4 days.

Genetic characterization. Single zoospore isolates were grown in antibiotic-amended V8 broth for 3 days at room temperature. Mycelial mats were washed, frozen, lyophilized, and ground with a sterile mortar and pestle. DNA was extracted with either a Qiagen Dneasy extraction kit (Qiagen, Valencia, CA) or via a cetyltrimethylammonium bromide (CTAB) procedure. A variety

of methods for generating molecular markers were tested for efficacy including isozyme, random amplified polymorphic DNA, and amplified fragment length polymorphism (AFLP). The AFLP technique resulted in a large number of reproducible markers and was chosen to characterize samples of P. capsici from Michigan. The AFLP technique involves cutting genomic DNA with moderately rare cutting (EcoRI) and frequent cutting (MseI) restriction enzymes, while concomitantly ligating synthetic adaptor fragments of DNA to the sticky ends created by the restriction enzymes (7). The result is a large number of DNA fragments that have ends with known DNA sequences. Amplification of fragment subsets (termed fingerprints) can be accomplished using polymerase chain reaction (PCR) primers complementary to the adaptor sequences with additional "selective" nucleotides. Changing the amount and type of selective nucleotides results in different subsets or fingerprints. Stringent PCR cycling parameters (touchdown technique) are used to ensure the fidelity of the reaction. For the analysis summarized here, adaptor sequences and fluorescent labeled selective primers were purchased as a kit through Perkin-Elmer ABI (Applied Biosystems, Foster City, CA). Using this system, AFLP fragments were resolved on a polyacrylamide gel by an ABI 377 gene sequencer. Fluorescent labels were excited by a laser and band emissions were analyzed in the form of an electropherogram where peaks represent individual bands. The sizing of fragments was particularly robust because a DNA ladder was loaded with every sample into the gel. To test for the reproducibility of fingerprints, DNA was extracted from a single isolate on three separate occasions approximately 3 months apart and subjected to the aforementioned protocol.

Data analysis. Isolates with identical multilocus AFLP finger-prints were considered to be members of the same clonal lineage and only a single representative was used for analysis. Because AFLP markers can only be scored confidently for presence (1) or absence (0), allele frequencies were estimated based on the assumption that populations under investigation meet the criterion for Hardy-Weinberg equilibrium, and that loci have only one "present" allele. The term population refers to all samples taken from a single field during a single year.

Genetic diversity within single populations was assessed by calculating the average number of polymorphic bands and estimating the average heterozygosity. Fixation indices were calculated according to methods of Weir and Cockerham (8) for populations from the same site over multiple years and among populations in Michigan using the program tools for population genetic analysis (TFPGA) (M. P. Miller, Northern Arizona University, Flagstaff). Confidence intervals for F statistics at the 95% confidence level were generated by bootstrapping at 1,000 iterations. The program NTSYS-pc version 2.02k (Exeter Software, Setauket, NY) was used to construct a similarity matrix from the presence/absence (1/0) data. Cluster analysis using the unweighted pair group with arithmetic averages (UPGMA) method was performed on the matrix and a tree was generated to give a visual representation of isolate similarity. Excoffier's ARLEQUIN program (L. Excoffier, University of Geneva) was used to assess population differentiation using a phenetic approach termed analysis of molecular variance (AMOVA), which allows for total genetic variation to be partitioned within and among populations using a classical analysis of variance (ANOVA).

#### RESULTS

Phenotypic results. Five isolates were recovered in 1997 from five different farms (four A1 and one A2 CT). One isolate was fully insensitive to mefenoxam, whereas the other four were fully sensitive. These findings prompted the extensive sampling conducted in 1998 in which 523 isolates (473 from cucurbits and 30 from bell pepper) were collected from 14 farms. A frequency histogram plotting percent growth of control on 100 ppm of

mefenoxam-amended media versus number of isolates revealed a trimodal distribution (3). Putative MS categories were assigned based on these groupings with sensitive (S) <30% growth of control, intermediately sensitive (IS) between 30 and 90% growth of control, and insensitive (I) >90% growth of control. In vitro crosses between isolates representative of the different putative sensitivity categories (S  $\times$  S, I  $\times$  S, IS  $\times$  S, and IS  $\times$  IS) resulted in progeny sets not significantly different than expected for insensitivity inherited as a single incompletely dominant gene unlinked to CT (P = 0.05) (3). In 1998, 55% of the isolates were sensitive to mefenoxam, 32% were intermediately sensitive, and 13% were fully insensitive to mefenoxam. A1 and A2 CTs were recovered in a ratio of approximately 1:1 in 8 of the 14 farms. Oospores were detected in naturally diseased cucurbit fruit from four farms, and 223 oospore progeny were recovered and germinated from a single diseased cucumber. All six possible MS × CT combinations were detected in this naturally occurring oospore progeny set (3).

In planta studies using sensitive, intermediately sensitive, and fully insensitive *P. capsici* isolates supported the in vitro screening categories, with sensitive isolates causing no disease on mefenoxam-treated plants, intermediately sensitive isolates being slowed by mefenoxam, and fully insensitive isolates showing no difference in the ability to colonize host tissue between treated and untreated plants at three times the field rate. All the progeny isolates were pathogenic on untreated pumpkin plants (K. H. Lamour and M. K. Hausbeck, *unpublished data*).

Sixty-three mefenoxam insensitive (18% intermediate and 82% fully insensitive) isolates were recovered from a single southwest Michigan field in 1998. Field experiments were conducted in this field during 1999 and 2000, testing alternative cultural control strategies, and no mefenoxam was applied. Two hundred isolates were recovered from this site over the course of the 1999 season and 34 isolates at the beginning of the 2000 season. Of the 200 isolates recovered in 1999 from this field, 141 had unique AFLP genotypes. Seventy percent of these were fully insensitive to mefenoxam, 28% were intermediately sensitive, and 2% were sensitive. In 2000, 15% of the isolates were intermediately sensitive and 85% were fully insensitive. A single fully insensitive clonal lineage rose in frequency over the course of the 1999 season and comprised 20% of the total number of samples recovered (4).

During 1999 and 2000, approximately 2,500 isolates were recovered from farms in Michigan. Both the A1 and A2 CTs were present in every field sampled, and mefenoxam insensitivity was detected in the majority of farms that had a history of mefenoxam use.

Genetic results. Nine populations from the four major vegetable production areas of Michigan were analyzed with the AFLP procedure (N = 641). AFLP analysis resolved a total of 94 clearly discernable markers when considering all the isolates together. No single isolate or group of isolates from a single location contained all 94 markers. The total number of AFLP loci in a single population ranged from 68 to 80. Seventeen (18%) fragments were fixed for the present state across all populations, 12 (13%) fragments were polymorphic in all populations, and 65 (69%) were fixed for presence or absence in some populations and polymorphic in others. The number of polymorphic bands within a single population ranged from 37 to 46 with estimated heterozygosities ranging from 0.18 to 0.22. Clonal reproduction was significant within single fields over the course of the growing season. For example, genotypic diversity in a single field ranged from 100% at the beginning of the growing season (seedling stage) to <30% at the time cucurbit fruit were ready for harvest (4). When considering all nine populations, genotypic diversity ranged from 42 to 96% with an average of 74% of the isolates in any sample set having unique genotypes. Although clonal reproduction was significant within single fields within years, no clones were recovered from single fields between years or among fields separated by at least 1 km. Fixation indices  $(\phi_{ST})$  between the populations sampled on consecutive years were very close to zero, indicating that gene diversity was not measurably impacted by genetic drift (5). The overall estimated  $\phi_{ST}$  for populations from different locations was 0.35, indicating that approximately 35% of the total genetic diversity present in Michigan *P. capsici* populations is found among populations and 65% is found within any one population. AMOVA partitioned genetic diversity among (40%) and within (60%) populations. The similarity tree based on UPGMA cluster analysis clearly showed that isolates from the

same site sampled over years branched from the same node, with no clustering of isolates based on the year of sampling. Cluster analysis also clearly showed that populations separated geographically branched from population-specific nodes (5).

#### DISCUSSION

During the past 10 years, Michigan has experienced a steady increase in the incidence of root, fruit, and crown rot on cucurbits

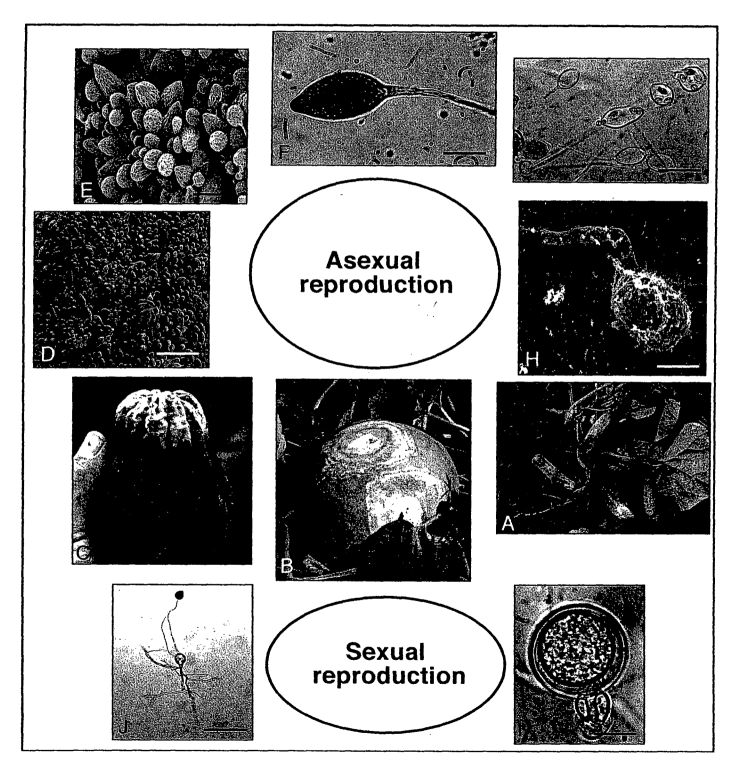


Fig. 1. Spore types and signs of infection caused by *Phytophthora capsici* on cucurbit fruit: A, infected cucumber, B, pumpkin, and C, acorn squash fruit. D, Scanning electron microscope (SEM) photo of an infected cucumber showing tufts of sporangia produced on the surface of the fruit ( $Bar = 300 \mu m$ ). E, Close-up of a single tuft of sporangia ( $Bar = 30 \mu m$ ). F, Typical papillate sporangium with a long pedicel ( $Bar = 20 \mu m$ ). G, Zoospores exiting sporangia after immersion in water ( $Bar = 50 \mu m$ ). H, SEM photo of a single encysted zoospore that germinated and directly penetrated the epidermis of a cucumber fruit ( $Bar = 4 \mu m$ ). I, Typical amphigynous oospore ( $Bar = 10 \mu m$ ). J, A germinating oospore with multiple germ tubes and a terminal sporangium ( $Bar = 100 \mu m$ ).

caused by *P. capsici*. Rotation to nonsusceptible hosts, in conjunction with cultural and chemical control strategies, have not provided economic control. Correspondence with other vegetable pathologists suggests that this phenomenon is not confined to Michigan, and a similar increase in control failures due to blight by *P. capsici* is being reported throughout the United States.

Investigation of the inheritance of MS demonstrated that MS is inherited as a single incompletely dominant gene unlinked to CT. In 1998, all six possible MS × CT combinations were present in single fields and insensitivity to mefenoxam was common in Michigan. Typical amphigynous oospores were observed in *P. capsici*-infected cucurbit fruit from multiple locations, and oospore progeny from a single naturally infected fruit showed segregation for MS and CT. These findings strongly support the hypothesis that sexual reproduction is occurring in the field, and also suggest that sexual recombination may directly generate progeny fully insensitive to mefenoxam. Tracking a single mefenoxam insensitive population over 2 years in the absence of mefenoxam selection pressure suggests that costs associated with mefenoxam insensitivity are minimal.

Estimates of average heterozygosity and polymorphism indicate surprisingly high levels of gene and genotypic diversity in all the populations of P. capsici analyzed. Tracking a single population through an entire growing season showed that asexual reproduction plays a significant role in disease development within a single season. Sampling single fields over consecutive years suggested that clones do not survive Michigan winters and that oospores are the primary survival propagule. Estimation of fixation indices for samples from the same site over consecutive years suggested that there was not a significant reduction in genetic diversity between growing seasons. This implies that populations are large enough to withstand dramatic effects of genetic drift. Cluster analysis revealed unambiguous groups corresponding to geographical locations with regional populations showing more similarity overall than populations from different regions. Population pairwise fixation indices corroborated this finding. The estimated overall fixation index and AMOVA are in agreement with both, suggesting that most (approx 60%) of the total genetic variability in Michigan is found within any one population, but that a relatively large component (40%) of genetic variability is found among populations.

Recommendations based on our findings are as follows: (i) the fungicide mefenoxam may be of limited usefulness because insensitivity appears to be selected for rapidly and is unlikely to decrease when mefenoxam selection pressure is removed; (ii) fields with epidemics are likely to harbor oospores for an extended amount of time (at least 5 years), and this factor must be considered before replanting to susceptible hosts; and (iii) factors that may contribute to the introduction of *P. capsici* into uninfested fields (e.g., drainage ditches between farms, irrigation ponds, and the dumping of culls) need to be considered and if possible avoided, because once an epidemic is established we have found no evidence that the population will become extinct in an agriculturally meaningful time period.

From an evolutionary perspective, it is clear that *P. capsici* has successfully colonized a number of geographical locations in

Michigan and that each of the populations sampled thus far have similarly high levels of genetic variability. The genetic stability of single populations over multiple years, the high fixation indices between even geographically close populations (1 km), and the clear structuring based on UPGMA cluster analysis all suggest that long-distance dispersal of inoculum is not common and that geographically isolated populations are also genetically isolated. It appears that the sexual stage of the P. capsici life cycle plays a significant role in survival as well as maintaining both genic and genotypic diversity, and has likely played a key role in the evolution of mefenoxam insensitivity. The combination of high levels of genetic variability, thick-walled oospores, and polycyclic asexual disease development make P. capsici a formidable pathogen (Fig. 1). This work underscores the need for management strategies aimed at preventing the spread of P. capsici to uninfested field sites and suggests that management strategies aimed at limiting spread within a single season may be the only option for growers with P. capsici-infested fields.

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#### Worksheet 4. Alternatives - Future Research Plans

Please describe future plans to test alternatives to methyl bromide. (All available methyl bromide alternatives from the alternatives list should have been tested or have future planned.) There is no need to complete a separate worksheet for future research plans for each alternative - you may use this worksheet to describe <u>all</u> future research plans.

1. Name of study:	Alternatives to methyl bromide for	control of Phytophthora capsici and Fusarium oxysporum	f.sp.
	melonis on cucurbits in Michigan.		
2. Researcher(s):	Dr. Mary Hausbeck		
	Mr. Brian Cortright		
3. Your test is plan	April to October, 2003,	, 2004	
4. Location:	Southwestern Michigan, USA at Michigan State University's Research and Extension Center,		,
	several plots will be placed with var	rious commercial growers.	
5. Name of alterna	tive to be tested:		
Multigard FFA (47, 7	71 gal/A)	Telone C-35 (15, 32 gal)	
Multigard Protect		Chloropicrin 100%	
Multigard Protect + \	√apam HL (37, 56 gal/A)	Idomethane 67/33	<del></del>
CX-100 (applied as	drip or preplant	Chicken manure composted	
6. Will crop yield b	e measured in the study?	YesXNo	
-		Whenever possible.	
alternatives have	e been tested and found unsu	plain why. (For example, the available uitable, an alternative has been identified but is rnatives are too expensive for this crop, etc.)	3
			<del></del>

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mide?	**************************************
	•
rota	tion, raised beds, b
	Approximate designations and dispersion in small class cash and approximate an
	Yes_X_ No
crobat	
	Yes No _X
ao on	
ge on	
	Yes No _X
4!	
sortium	
g in	·
	\$ 1.1 million
vi bromi	ide. Describe each
yı bi olili	de. Describe each
earch its t	op priority.
	·
of meth	ıyl bromide
	pital outiay).
of new di	sease management strategies
<del></del>	
··	

#### Worksheet 5. Additional Information

1.	How will you minimize your u	se and/or emissions of methyl bromi	de?
	1a. Check all methods you will use	X Tarpaulin (high density polyethylene)  X Virtually impermeable film (VIF)  X Cultural practices (please specify)	rotation, raised beds, b
	1b. Will you use other pesticides to r	trickle irrigation educe use of methyl bromide?	Yes_X_ No
	If yes please specify. <u>fo</u>	liar fungicides including Ridomil Gold and Acro	bat
	1c. Other non-chemical methods: (pl raised beds, trickle, black plastic, folia		
2.	Do you have access to recycle	ed methyl bromide?	Yes No X_
	If yes, how many pounds?	lbs.	,
3.	Do you anticipate that you wil January 1, 2005?	I have any methyl bromide in storage	e <b>on</b> Yes No X
	If yes, how many pounds?	lbs.	
	on research to develop alternation 1992)?	nt spent to date by the user or consor atives to methyl bromide (beginning i de to reduce your reliance on methyl	n \$ 1.1 million
		plant pathology program has made the resear	cah ita tan priority
	Michigan State Offiversity's vegetable	plant pathology program has made the resear	on its top priority.
6.	_	ow you to stop or reduce your use of pesticide; completion of research pla	•
	Completion of our research plan, iden	tification, development, and implementation of	new disease management strategies
	would greatly reduce our methyl brom	ide use.	
	When do you expect these to oc	cur? Within 5 to 10 years.	
7.	Range of acres farmed by grow (insert number of users in each of	wers included in this application?	
	<b>4</b> 0-10 acres		
	<b>8</b> 10-25 acres		•
	<b>6</b> 25-50 acres		
	<b>5</b> 50-100 acres	•	
	6 100-200 acres		
	<b>1</b> 200-400 acres		
	over 400 acres		

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#### Worksheet 5. Additional Information (continued)

this application will apply methyl bromide? (insert no in each category)	s included in umber of users
0 - 5,000 sq. ft. 5,001 - 10,000 sq. ft.	
10,001 - 20,000 sq. ft.	
20,001 - 40,000 sq. ft. 40,001 - 80,000 sq. ft.	
4 80,001 - 160,000 sq. ft.	
25 over 160,000 sq. ft.	
I certify that all information contained in this document is factual to	o the best of my knowledge.
Signature Mary K. Hausbeck	Date
Print Name Mary K. Hausbeck	Title Professor
	· /
Information in this application may be aggregated with information States government to justify claims in the national nomination pactonsidered "critical" and authorized for an exemption beyond the crucial to making compelling arguments in favor of critical use exeassert any claim of confidentiality that would affect the disclosure information contained in this application.	ckage that a particular use of methyl bromide be 2005 phaseout. Use of aggregate data will be emptions. <b>By signing below</b> , you agree not to
States government to justify claims in the national nomination pactonsidered "critical" and authorized for an exemption beyond the crucial to making compelling arguments in favor of critical use exercises assert any claim of confidentiality that would affect the disclosure	ckage that a particular use of methyl bromide be 2005 phaseout. Use of aggregate data will be emptions. <b>By signing below</b> , you agree not to

Burden means the total time, effort, or financial resources expended by persons to generate, maintain, retain, or disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; develop, acquire, install, and utilize technology and systems for the purposes of collecting, validating, and verifying information, processing and maintaining information, and disclosing and providing information; adjust the existing ways to comply with any previously applicable instructions and requirements; train personnel to be able to respond to a collection of information; search data sources; complete and review the collection of information; and transmit or otherwise disclose the information. Public reporting burden for this collection of information is estimated to average 324 hours per response and assumes a large portion of applications will be submitted by consortia on behalf of many individual users of methyl bromide. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a current OMB control number.

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Worksheet	6 Ann	lication	Summary

This worksheet will be posted on the	e web to notify the public of requests for critical use exemptions beyond the 2005 phase out for methyl bromide. Ther	efore, this worksheet cannot be claimed a
1. Name of Applicant:	Michigan cucurbit growers	
2. Location:	Michigan, USA	•
3. Crop:	Cucurbits including: watermelon; muskmelon, cucumber, summer squash, winter squash	
4. Pounds of Methyl Bromide Requ	uested 2005 62,142 a.i.	
5. Area Treated with Methyl Bromio	de 2005 1,446 acres units	
6. If methy! bromide is requested for	or additional years, reason for request:	
Additional time is needed to develop	effective alternatives for Phytophthora capsicl. Michigan State University has an active research program, and is making progress in diseas	e management.
	A Total AMO and the	20 miles - 10 miles -
2006 60,970 lbs		
200758,625 ibs	Area Treated 1,364 acres units	

Place an "X" in the column(s) labeled "Not Technically Feasible" and/or "Not Economically Feasible" where appropriate. Use the "Reasons" column to describe whaternative is not feasible.

Not Technically Feasible	Not Economically Feasible	Reasons
х		Not effective,
×		Not effective.
×		Not effective, pathogens long-lived.
x	-	Efficacy is not proven, requires solarization.
×		Climate in Michigan, USA is too cold.
x		Not technically feasible for large scale agriculture.
x		Efficacy is not proven.
×		Not effective, already used in commercial production.
×		Not effective, pathogens long-lived, already used in commercial production.
x		Efficacy is not proven.
x		Flooding is not feasible, trickle and reised beds are used, but frequent heavy rains favor disease.
×		Utilized by growers, but is not adequate for disease control.
x		Resistant rootstock has not been identified. Would not be effective against root rot.
×		Not effective, many growers already using techniques.
x		Resistant varieties have not been identified.
x		Volcanic ash, rockwool are not viable alternatives for large-scale production in Michigan, USA.
x		Primary pathogens are not disseminated on seed or transplants.
	Technically Feasible  X  X  X  X  X  X  X  X  X  X  X  X  X	Technically Feasible  X  X  X  X  X  X  X  X  X  X  X  X  X